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## Peyman

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## (54) METHOD AND COMPOSITION FOR HYPERTHERMALLY TREATING CELLS

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	A61K 9/51	(2006.01)
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	A61N 7/02	(2006.01)
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	B82Y 99/00	(2011.01)
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(2006.01)

U.S. Cl. CPC ...... A61M 37/0092 (2013.01); A61B 3/13 (2013.01); A61B 5/0095 (2013.01); A61B 5/4836 (2013.01); A61B 6/032 (2013.01); A61B 6/037 (2013.01); A61B 18/04 (2013.01); A61F 7/00 (2013.01); A61F 9/00 (2013.01); A61K 9/0004 (2013.01); A61K 9/0009 (2013.01); A61K 9/5169 (2013.01); A61K 31/713 (2013.01); A61K 35/30 (2013.01); A61K 38/185 (2013.01); A61K 39/12 (2013.01); A61K 41/0052 (2013.01); A61K 45/06 (2013.01); A61K 47/48569 (2013.01); A61K 47/48861 (2013.01); A61K 47/48884 (2013.01); A61K 48/0041 (2013.01); A61K 49/227 (2013.01); A61M 5/007 (2013.01); A61N 2/002 (2013.01); A61N 5/062 (2013.01); A61N 5/1001 (2013.01); A61N 5/1017 (2013.01); C12N 7/00 (2013.01); A61B 5/01 (2013.01); A61B 8/0841 (2013.01); A61B 8/10 (2013.01); A61B 8/4416 (2013.01); A61B 18/12 (2013.01); A61B 18/18 (2013.01); A61B 2017/00084 (2013.01); A61B 2017/00106 (2013.01); A61B 2018/00791 (2013.01); A61B 2090/378 (2016.02); A61F 9/0079 (2013.01); A61F 2007/0004 (2013.01); A61F 2009/00863 (2013.01); A61M 2037/0007 (2013.01); A61N 2/02 (2013.01); A61N 5/025 (2013.01); A61N 5/0625 (2013.01); A61N 7/02 (2013.01); A61N 2005/0662 (2013.01); A61N 2005/1098 (2013.01); B82Y 99/00 (2013.01); C12N 2700/00 (2013.01); G01R 33/4804 (2013.01); G01R 33/4808 (2013.01)

## (58) Field of Classification Search

None

See application file for complete search history.

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Primary Examiner — Yunsoo Kim (74) Attorney, Agent, or Firm — Thompson Hine LLP

## (57) ABSTRACT

A method and composition for hyperthermally treating tumor cells in a patient under conditions that affect tumor stem cells and tumor cells.

### 23 Claims, 6 Drawing Sheets

## Related U.S. Application Data

continuation-in-part of application No. 14/679,083, filed on Feb. 17, 2015, now Pat. No. 9,320,813, which is a continuation-in-part of application No. 14/624,334, filed on Feb. 17, 2015, now Pat. No. 9,302,087, which is a continuation-in-part of application No. 14/554,840, filed on Nov. 26, 2014, Pat. No. 9,017,729, which is a continuation-in-part of application No. 14/311,464, filed on Jun. 23, 2014, now Pat. No. 8,932,636, which is a continuation-in-part of application No. 13/915,282, filed on Jun. 11, 2013, now Pat. No. 8,668,935, which is a continuation-in-part of application No. 13/665,458, filed on Oct. 31, 2012, Pat. No. 8,709,488, which is a continuation-in-part of application No. 13/610,503, filed on Sep. 11, 2012, now Pat. No. 8,795,251, which is a continuation-in-part of application No. 13/527,005, filed on Jun. 19, 2012, now Pat. No. 8,808,268, which is a continuation-in-part of application No. 13/455,237, filed on Apr. 25, 2012, Pat. No. 8,801,690, which is a continuation-in-part of application No. 13/361,786, filed on Jan. 30, 2012, now Pat. No. 8,481,082, which is a continuation-in-part of application No. 13/307,916, filed on Nov. 30, 2011, now Pat. No. 8,119,165, which is a continuation-in-part of application No. 13/189,606, filed on Jul. 25, 2011, now Pat. No. 8,137,698, which is a continuation-in-part of application No. 13/149,209, filed on May 31, 2011, 7,964,214, now Pat. No. which continuation-in-part of application No. 12/478,029, filed on Jun. 4, 2009, now Pat. No. 7,638,139, which is a continuation-in-part of application No. 11/485,352, filed on Jul. 13, 2006, now Pat. No. 7,101,571.

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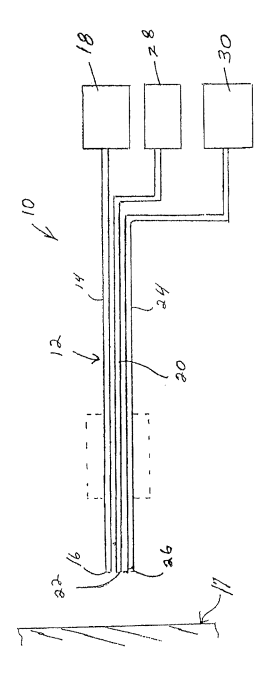
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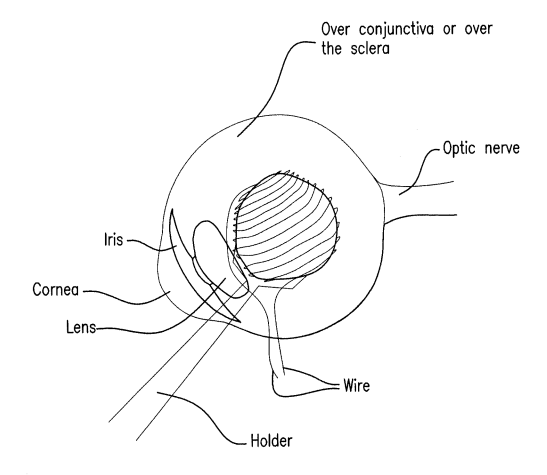


FIG.2

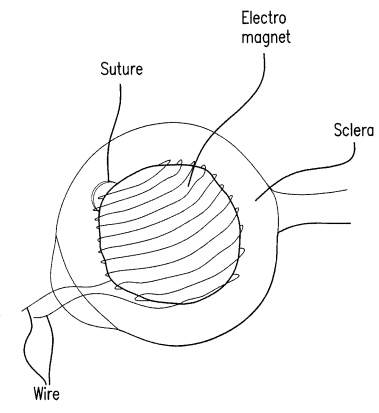


FIG.3

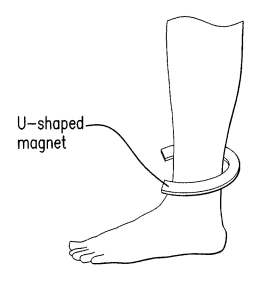


FIG.4A

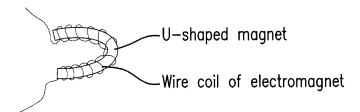
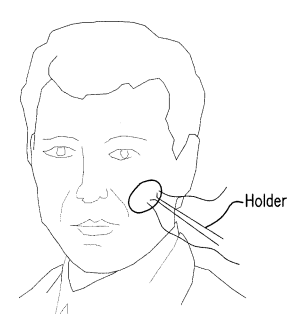


FIG.4B

Jul. 19, 2016



The electromagnet can be converted also to an alternating magnet.

All patients can be treated with an alternating magnet, electromagnetic radiation or ultrasound.

FIG.5

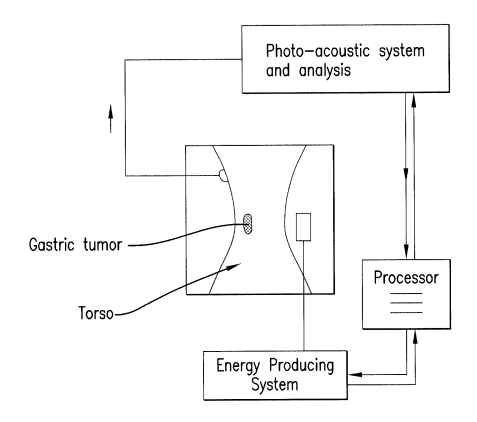


FIG.6

# METHOD AND COMPOSITION FOR HYPERTHERMALLY TREATING CELLS

This application is a Continuation-In-Part of U.S. patent application Ser. No. 14/724,577 filed May 28, 2015 now U.S. Pat. No. 9,320,813; which is a Continuation-In-Part of U.S. patent application Ser. No. 14/679,083 filed Apr. 6, 2015 now U.S. Pat. No. 9,302,087; which is a Continuation-In-Part of U.S. patent application Ser. No. 14/624,334 filed Feb. 17, 2015 now U.S. Pat. No. 9,289,491; which is a Continuation-In-Part of U.S. Ser. No. 14/554,840 filed Nov. 26, 2014 now U.S. Pat. No. 9,233,158; which is a Continuation-in-Part of U.S. patent application Ser. No. 14/311,464 filed Jun. 23, 2014 now U.S. Pat. No. 9,017,729; which is a Continuationin-Part of U.S. patent application Ser. No. 13/915,282 filed Jun. 11, 2013 now U.S. Pat. No. 8,932,636; which is Continuation-in-Part of U.S. patent application Ser. No. 13/665, 458 filed Oct. 31, 2012 now U.S. Pat. No. 8,668,935; which is a Continuation-in-Part of U.S. patent application Ser. No 13/610,503 filed Sep. 11, 2012 now U.S. Pat. No. 8,709,488; <sup>20</sup> which is a Continuation-in-Part of U.S. patent application Ser. No. 13/527,005 filed Jun. 19, 2012 now U.S. Pat. No. 8,795,251; which is a Continuation-in-Part of U.S. patent application Ser. No. 13/455,237 filed Apr. 25, 2012 now U.S. Pat. No. 8,808,268; which is a Continuation-in-Part of U.S. 25 patent application Ser. No. 13/361,786 filed Jan. 30, 2012 now U.S. Pat. No. 8,801,690; which is a Continuation-in-Part of U.S. patent application Ser. No. 13/307,916 filed Nov. 30, 2011 now U.S. Pat. No. 8,481,082; which is a Continuationin-Part of U.S. patent application Ser. No. 13/189,606 filed 30 Jul. 25, 2011 now U.S. Pat. No. 8,119,165; which is a Continuation-in-Part of U.S. patent application Ser. No. 13/149, 209 filed May. 31, 2011 now U.S. Pat No. 8,137,698; which is a Continuation-in-Part of U.S. patent application Ser. No. 12/478,029 filed Jun. 4, 2009 now U.S. Pat. No. 7,964,214; 35 which is a Continuation-in-Part of U.S. patent application Ser No. 11/485,352 filed Jul. 13, 2006 now U.S. Pat. No. 7,638, 139; which is a Division of U.S. patent application Ser. No. 10/073,863 filed Feb. 14, 2002 now U.S. Pat. No. 7,101,571; the entirety of each is hereby expressly incorporated by ref-  $^{40}$ erence herein.

## FIELD OF THE INVENTION

The present invention relates to a method and composition 45 for hyperthermally treating cells at a site in the body. More particularly, the present invention relates to a method for treating cells at a target site in the body, such as at a lens capsule of an eye, tumors, and exudative or wet age related macular degeneration (AMD) by applying thermal energy to 50 the target site to heat the cells to a temperature which will kill the cells or impede cell multiplication without exceeding the protein denaturation temperature of the tissue.

## BACKGROUND OF THE INVENTION

Several techniques currently exist for treating cells at a selected site in the body with heat or chemicals to kill or impede multiplication of those cells to prevent undesired cell proliferation. For example, numerous types of chemotherapy 60 drugs exists which, when injected into a tumor or delivered systemically to a patient, attack and kill cancerous cells to prevent them from further multiplying. However, unless the treatment affects the tumor stem cells that have mutated to result in uncontrolled tumor growth and metastasis, treatment 65 will not be effective. Stem cells are pluripotential undifferentiated cells. Tumor stem cells are frequently located in the

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bulk tumor mass, are involved in tumor metastasis, and often elude detection. Over time, stem cells become resistant to standard chemotherapy regimens by constant genetic mutations that confer resistance.

Thermal radiation techniques can also be used to kill cancerous or other undesired cells. Cell death begins to occur when the cells are heated to a temperature of about 5° C. or more above the normal body temperature of 37° C. Applying thermal radiation to a localized site in the body, such as a tumor or other area containing undesired cells, can heat the cells at the site to temperatures in excess of 60° C. Such high temperatures causes a phenomenon known as protein denaturation to occur in the cells, which results in immediate cell death. Accordingly, thermal radiation therapy has been suitable in successfully treating certain types of cancers and other diseases involving uncontrolled cell growth.

Other types of heating techniques, such as the use of probes or catheters to provide localized heat to a site of interest also exist. Like thermal radiation therapy, these techniques also heat the cells to a temperature sufficient to cause protein denaturation in the cells to thus kill the cells quickly.

Photosensitive chemicals are also used to kill cells at certain sites of interest in the body. For example, a photosensitive chemical can be injected directly into a site of interest to expose cells at that site to the chemical. A light emitting source, which emits light at a wavelength that will activate the photosensitive chemical, is then focused on the site of interest. Accordingly, the light activates the photosensitive chemical that has been absorbed by or is otherwise present in the cells of interest. The activated chemical kills the cells, which thus prevents undesired cell proliferation.

Although the techniques mentioned above can be suitable for preventing certain types of cell proliferation at certain sites in the body, several drawbacks with these techniques exist. For example, often the use of chemotherapy drugs alone to treat a tumor or cancerous site is insufficient to kill the undesired cells. The only current treatments directed at tumor stem cells are heavy doses of ionizing radiation, thermal radiation or thermotherapy, or chemotherapy. Moreover, the chemotherapy drugs and other treatments also indiscriminately kill many normal healthy cells along with the cancerous cells, which can adversely affect the patient's health and are frequently ineffective against late stage cancers.

The use of ionizing radiation in conjunction with chemotherapy can have a more detrimental effect on the cancerous cells. However, as with chemotherapy, ionizing radiation often kills normal healthy cells, such as those in front of or behind the site of interest, along with the cancerous cells. Moreover, the intense heating of the cells can cause the cells to coagulate and thus block the capillaries at the site of interest. The blocked capillaries therefore prevent chemotherapy drugs from reaching the site of interest.

One example of a method of chemically treating a target site is disclosed in U.S. Pat. No. 6,248,727 to Zeimer. This method delivers a liposome containing a fluorescent dye and tissue-reactive agent. The liposome is administered intravenously to flow to the locus in the eye of the patient and the site is non-invasively heated to release the dye and the tissue-reactive agent. The dye is fluoresced to observe the pattern of the fluorescence. The tissue-reactive agent is activated to chemically damage and occlude the blood vessel. The liposomes are selected to release the dye at a temperature of 41° C. or less without causing thermal damage to the blood vessel.

In addition, the above techniques have not been used to prevent unwanted cell proliferation at certain locations in the eye, such as at the retina or at the lens capsule. Because the

retina is very sensitive, conventional ionizing radiation techniques can be too severe to treat cancerous cells on, in or under the retina.

Also, after cataract surgery, a phenomenon known as capsular opacification and, in particular, posterior capsular 5 opacification can occur in which the epithelial cells on the lens capsule of the eye experience proliferated growth. This growth can result in the cells covering all or a substantial portion of the front and rear surfaces of the lens capsule, which can cause the lens capsule to become cloudy and thus 10 adversely affect the patient's vision. These cells can be removed by known techniques, such as by scraping away the epithelial cells. However, it is often difficult to remove all of the unwanted cells. Hence, after time, the unwanted cells typically will grow back, thus requiring further surgery.

Accordingly, a need exists for a method for hyperthermally treating tissue and preventing unwanted cell proliferation at sites in the body, especially at sites in the eye such as the retina, choroid and lens capsule, which does not suffer from the drawbacks associated with the known techniques dis- 20 cussed above. The method should also treat tumor stem cells to target and eradicate a tumor source and eliminate or slow tumor metastasis. The method should also target and damage the specific tumor-associated vasculature, in effect, starving the tumor of its nutrient supply.

## SUMMARY OF THE INVENTION

The present invention is directed to a method of hyperthermally treating tissue by heating the tissue above a temperature 30 which kills cells in the tissue. In particular, the invention is directed to a method of heating tissue above a temperature effective to treat the tissue without denaturing the protein. The present invention also relates to a method and composition for hyperthermally treating cells in the eye with simul- 35 taneous imaging.

One embodiment of the method targets tumor stem cells by combining chemotherapy with thermotherapy to target tumor stem cells while leaving normal cells either unaffected or minimally effected by the therapy. This embodiment of the 40 heating and detecting a temperature of a tissue between a first method uses increases in temperature to first prime the tumor stem cells, and then to kill the tumor stem cells synergistically with thermotherapy. In one embodiment, nanoparticles that are coated with or otherwise contain antibodies that specifically target certain cells and/or cell types, e.g., tumor cells. 45 This helps to minimize or eliminate chemotherapy adverse effects on non-tumor bearing vital organs such as liver, kidney, gut, heart, central nervous system, while providing supra-therapeutic doses of the chemotherapy drugs to tumor cells, both local and metastatic, and stem cells. In this way, the 50 duration of chemotherapy administered to a patient may be reduced without compromising effective treatment.

One embodiment of the method targets tumor stem cells by combining localized internal ionizing radiation therapy with thermotherapy to target tumor stem cells and to destroy their 55 associated vasculature, while leaving normal cells and nontumor vasculature either unaffected or minimally affected by the therapy. This embodiment of the method uses increases in temperature to first prime the tumor stem cells, and then to kill the tumor stem cells synergistically with thermotherapy, and 60 uses localized internal ionizing radiation therapy to destroy vasculature associated with the tumor. In one embodiment, nanoparticles that are coated with or otherwise contain antibodies that specifically target certain cells and/or cell types, e.g., tumor cells, also contain a radioactive isotope. The 65 increased temperature from the thermotherapy kills the tumor stem cells, both local and metastatic. The localized internally

administered radiotherapy damages the proximate and/or adjacent endothelial cells of the vasculature feeding the tumor cells, both local and metastatic. The embodiment of the method may be used in a patient receiving anti-vascular endothelial growth-factor therapy to provide an additional source of vessel damage. This combination therapy increases the likelihood of eradication of tumor cells and stem cells associated with the tumor by additionally damaging or obliterating the vessels that provide their nutrient supply. The result is more robust therapy and increased chances for patient survival.

Accordingly, a primary aspect of the invention is to provide a method for heating tissue at least to a temperature sufficient to hyperthermally treat the tissue.

Another aspect of the invention is to provide a method of hyperthermally treating tissue to a temperature sufficient to kill cells in the tissue and at a temperature below the protein denaturization temperature of the tissue.

A further aspect of the invention is to provide a method of hyperthermally treating tissue, where the tissue includes or is provided with a temperature indicator to indicate a hyperthermally effective temperature of the tissue.

Still another aspect of the invention is to provide a method 25 of hyperthermally treating tissue where a temperature indicator composition is introduced into the tissue or bloodstream near the tissue to indicate a tissue temperature effective to hyperthermally treat the tissue and a temperature indicator to indicate a tissue temperature above a protein denaturization temperature of said tissue.

A further aspect of the invention is to provide a method of hyperthermally treating tissue by introducing a temperature indicator into the tissue and heating the tissue to a temperature where the temperature indicator can be detected. In a preferred embodiment, the temperature at which the indicator can be detected is a temperature effective to hyperthermally treat the tissue and is at a temperature below the protein denaturization temperature.

A further aspect of the invention is to provide a method of temperature and a second temperature. The method introduces a temperature indicator into the tissue. The temperature indicator includes a first dye that can be detected at the first temperature to indicate that the first temperature has been reached, and a second dye that can be detected at the second temperature to indicate that the second temperature has been reached.

Still another aspect of the invention is to provide a temperature indicating composition for introducing into a tissue to be thermally treated. The composition includes a first dye encapsulated in a heat sensitive liposome where the first dye is releasable at a temperature effective to hyperthermally treat the tissue and at a temperature below the protein denaturization temperature. The composition also includes a second dye encapsulated in a second liposome where the second dye is releasable at a temperature at or above the protein denaturization temperature.

Another aspect of the invention is to provide a method to hyperthermally treat tissue to kill the tissue cells substantially without protein denaturization of the tissue where the tissue includes a heat sensitive liposome containing a temperature indicating dye and a temperature activated bioactive compound. The tissue is heated to release the dye from the liposome to indicate a thermally effective temperature to kill cells in the tissue at a temperature below the protein denaturization temperature. The heat applied to the tissue simultaneously releases the bioactive compound to treat the tissue.

Another aspect of the invention is to provide labeled nanoparticles, either alone or conjugated with activatable cellpenetrating peptides (ACPPs) to assist in localizing and destroying cancer cells or tumor tissue remaining after surgical excision of a tumor. Another aspect of the invention relates 5 to ACPP-conjugated nanoparticles that can also be adapted to attach to normal cell specific receptors in order to stain and treat the tissues. These cells can be as involved with vasculature such as myofibroblasts becoming activated after stent surgery; or nerve cells that create neuromas. Another aspect of the invention relates to the treatment or debulking of other unwanted masses, such as fibromas, meningiomas, adenomas, or degenerative cells causing Parkinson's Disease or of nerves involved in transmitting pain, e.g., for patients in chronic pain. Another aspect of the invention is to treat malignant tumors, such as neuroblastoma, melanoma, skin, breast, brain kidney, lung, intestinal, and genitourinary, bone, gland, and blood cancers. Another aspect of this invention is to target normal cells of the body responsible for an immune response, 20 such as lymphocytes, etc., that prevent an organ transplant. Another aspect of the invention is to treat undesirable vascular rupture and bleeding.

Another aspect of the invention is to target and damage tumor cells, tumor stem cells, and endothelial cells of tumor- 25 associated vessels, e.g., blood vessels, lymphatics.

Another embodiment of the invention is to provide immunotherapy in combination with thermal therapy to a target site. Thermotherapy is provided by administering a composition comprising a targeting agent and/or antitumor-antibody-labeled nanoparticle where the nanoparticle forms a targeting agent and/or antibody labeled nanoparticle-cell complex at a tumor site, exposing the patient to an energy source where the nanoparticles respond by

source where the nanoparticles respond by expanding thus generating an acoustic signal, measuring the acoustic signal and relating the measured signal to a temperature of the nanoparticle-cell complex, and controlling the temperature of the nanoparticle-cell complex from about 39° C. to about 58° C. based on the acoustic signal to hyperthermally damage or kill cells in the tumor; and providing immunotherapy by providing an immunotherapy agent associated with the nanoparticles, providing an immunotherapy agent not associated with the nanoparticles, and/or providing a therapeutic concentration of autologous leukocytes to the 45 patient, where immunotherapy enhances a patient's immune response. In specific embodiments, thermotherapy is performed in the absence of a contrast agent, substantially simultaneously with, prior to, or subsequent to immune therapy. The immunotherapy may reduce immunosuppression. The thermotherapy may supplement chemotherapy and/or radiation therapy. A marker may be evaluated to assess a therapy effect. The nanoparticle may be associated with a member of specific binding pair such as streptavidin-biotin pair, a cellular receptor-agonist or antagonist pair, an enzyme-substrate pair, and/or an antibody-antigen pair. The nanoparticle may be associated with a virus, e.g., a modified virus, a tumoricidal virus, and/or an adeno-associated virus (AAV) or virus like particles (VLP) made from modified plant viruses. VLP, plant viral capsids or membranes lacking the genetic components of the virus and thus unable to multiply in the organism, are specifically used with immune therapy to enhance a cellular immune response in the body towards specific antigen such as the tumor cells. The use of VLP eliminates the need for adjuvants in immune therapy for enhancing the immune response to the tumor cells.

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In one embodiment, hyperthermal therapy is used for gene delivery. In one embodiment, hyperthermal therapy is used for gene therapy. In one embodiment, hyperthermal therapy is used for medication delivery.

For gene delivery, the use of hyperthermal therapy provides a method to achieve precise, accurate, and reproducible control of genes provided to the patient in nanoparticles, including quantum dots. For example, in such use, a goal may be to selectively provide a gene, contained or carried in nanoparticles or quantum dots, to malignant cells, while not providing the gene to non-malignant (i.e., normal) cells. The use of hyperthermal therapy permits delivery and/or activation of the gene only to the selective location, using temperature as a ready and precise regulatory element. The skilled person in the art will appreciate that the nanoparticle structure itself may contain, be coated with, and/or otherwise comprise compounds that have temperature-dependent sensitivity and properties to achieve further refinement and sensitivity of the inventive method. The result is exquisite sensitivity to reducing or eradicating cancer cells, and sparing of normal nonmalignant cells, in such a cancer therapy approach.

It is known that one type of cancer therapy involves gene therapy. However, it is also known that gene therapy is not limited to cancer treatment, and has wide applicability, with exemplary but non-limiting uses in, e.g., ocular therapy to treat age-related macular degeneration and diabetic retinopathy; neural therapy to treat Parkinson's disease and Alzheimer's disease; etc. For such gene therapy, the use of hyperthermal treatment is important in treating such pathological conditions. Hyperthermal therapy can be accurately and reproducibly delivered, controlled, and monitored, rendering it an improvement in both ease of delivery and predictability compared to other methods known in the art such as, e.g., electroporation, nanoparticle delivery, quantum dot delivery, a modified viral carrier (vector), etc. Hyperthermal delivery with nanoparticles specifically provides a gene or genes or cell delivery, even with a viral carrier, to specific cells to which the antibody and chitosan-coated nanoparticles attach. At about 42° C., a temperature created by the thermal energy unit under the control of the photoacoustic system, the genes along with CRISPR/Cas or CRISPER/Cpf1 are released from the thermosensitive polymers, such as chitosans, and migrate in the cell.

Administration of hyperthermal therapy thus benefits both the patient and the medical provider. Hyperthermal therapy can additionally be used for overall, general, or en masse treatment of any organ without limitation, or a specific region of such an organ, including but not limited to brain, eye, breast, liver, pancreas, bone, heart, kidney, bladder, male or female reproductive organs, nerves, etc. In one embodiment, the method is used in vitro, e.g., in cultured cells. In one embodiment, the method is used in vivo, e.g., in a patient.

When the inventive method is used for cancer therapy, one uses one or more orders of magnitude higher numbers of nanoparticles than what is needed for gene delivery.

Genes, once in the nucleus, undergo transcription and translation into the specific proteins or protein channels (Narayanan et al., Scientific Reports 3, article number 2184, doi: 10.1039/srep02184, describing a mimic of microtubule mediated protein transport using designed biotinylated peptides with microtubule-associated sequences (MTAS) and a nuclear localization signaling (NLS) sequence, conjugating with streptavidin-coated CdSe/ZnS quantum dots to enhance endosomal escape and promote targeted nuclear delivery into mesenchymal stem cells by microtubules; Ho et al., J. Vis Exp. 30 (2009) 1432, describing a specific quantum dotlabeled DNA complex (Cy5) that forms by electrostatic self-

assembly, facilitating DNA cellular update and protecting against DNA degradation and monitored using combined quantum dot-FRET and microfluidics).

In any of these embodiments previously described, the method advantageously reduces the time for therapy to less than one second for gene therapy. A burst of thermal energy under the control of photoacoustic technology is sufficient to heat the nanoparticles to 60° C. or more for a very short time, e.g., one femtosecond ( $10^{-15}$  second) to one second to a few minutes. When in contact with the cell membrane, the heated nanoparticles rupture, all at once, all cell membranes in contact with or proximate to the nanoparticles. The heated nanoparticles damage the cell membrane and create holes or pores permitting the gene(s) or medication to freely enter into cells.

This temperature increase provided by the hyperthermal 15 therapy has a two-fold benefit. It results in release of gene(s) and/or medications that are attached to the nanoparticles or that are present in the vicinity of the cells. It facilitates entry of the gene(s) and/or medications into the cell interior, to create very small openings or holes; i.e., the localized thermal energy is sufficient to make holes in the cell membrane. Hyperthermal conditions in the presence of nanoparticles and certain toxins create holes or pores in the cellular membrane of cancer cells. Typically, normal healthy cell 25 membranes are comprised of the phospholipids phosphatidylserine (PS) and phosphatidylethanolamine (PE), which are located within the inner membrane facing the cell interior. However, for cancer cells, PS and PE face the outer membrane surface, they are "flipped" in cancer cells (Leite et al. 30 Biophysical Journal 109 (2015) 936 which is expressly incorporated by reference herein in its entirety). The melting point PS and PE is about 65° C.-70° C. When the nanoparticles are heated to 65° C.-70° C., the exposed lipids of the cell membrane melt and create a dehesence in the cell membrane, 35 through which material such as protein, RNA or DNA etc. move inside the cells.

In the inventive method, the temperature for hyperthermal therapy is controlled with photoacoustic technology during therapy, and is maintained at the desired level for the defined 40 time. Coating the nanoparticles with phospholipase and antiphospholipid antibodies and toxins, such as snake, scorpion, and bee venom, etc., can enhance the lysis effect of the antibody-coated nanoparticles to the cell membrane of the cancer cell, thus enhancing the hyperthermal effect.

These thermal energy-generated holes in the cell membrane simplify entry of material to the interior of the cell. In cancer therapy, the goal is either to kill a cancer cell, or to silence certain genes in the cancer cell and deliver the medication, while the organ structure or a tumor and the temperature is imaged and thermal energy is controlled. In an embodiment where the targeted cells is a cancer cell, the cancer cells are damaged or killed with medications now internalized into the cancer cell through the small holes in the cell membrane. This method also modifies the cancer cell genes in a very 55 short period of time. In an embodiment where the targeted cell is a normal cell, the same method is used for en masse gene delivery to an organ, such as the retina, or even to cells in vitro. The desired gene or genes replace the defective gene or genes and imaging of the organ and temperature control occur 60 simultaneously. This method is beneficial compared to existing techniques using vectors or to nanoparticle carriers alone. For example, vectors do not provide sufficient yields of transfection, and also can produce side effects of viral delivery. In contrast, the disclosed method delivers with ease antibody 65 coated nanoparticles conjugated with the appropriate genes along with the CRISPR complex to the cells en masse in a

short period of time. Desired cells are transfected en masse by modifying their genes or delivering new genes to the nucleus or the mitochondria. the delivered genes or protein can be protected by a wrapper of single stranded DNA which can tolerate thermal energy.

As previously disclosed, the inventive method can modify a defective gene in a cell, i.e., the method is not limited to therapy of malignant cells although the skilled person in the art appreciates that a defective gene may be a sole or contributing factor to a malignancy.

Such genes include, but are not limited to, genes involved in cell membrane polarization and depolarization, genes encoding opsin family genes, genes encoding neurotransmitters, genes encoding regulatory proteins, genes encoding products for synthesis or inhibition of a therapeutic protein, tumor suppressor genes, therapeutic genes, defective genes, oncogene suppressor genes, genes that have an stimulatory action in response to light or ultrasound, etc.

A gene or genes involved in cell membrane polarization accomplished by damaging the cellular membrane sufficient 20 and depolarization can create an action potential in membrane of both excitable and non-excitable cells, and also of tumor cells. In the case of tumors, their cell membranes can be rendered permanently depolarized, beneficially facilitating access to desired agents that induce cell damage and death, e.g., anti-cancer agents, more specifically, alkylating agents; platinum coordination agents such as cisplatin, carboplatin, and oxaliplatin; antimetabolites; microtubule damaging agents such as paclitaxel and vincristine; topoisomerase inhibitors such as etoposide; antibiotics such as mitomycin C, bleomycins, and doxorubicin; and agents such as hydroxyurea and L-asparaginase. Such genes include, but are not limited to, genes producing a stimulatory action in response to, e.g., light, ultrasound, etc. such as genes encoding opsin family members that include rhodopsin, photopsins, halorhodopsin, channelrhodopsins, etc., genes encoding neurotransmitters such as genes encoding glutamate/aspartate transporters, GABA transporters, glycine transporters, monoamine transporters such as dopamine transporters, norepinephrine transporters, serotonin transporters, and vesicular monoamine transporters. Such gene or genes can be simulated with light to effect therapy in excitable cells such as the retina, brain, or heart. Alternatively, such gene or genes can modify a genetic defect of these cells.

In one embodiment, adding an opsin gene renders any cell 45 membrane, not limited to membranes of naturally excitable cells such as retina, brain, and cardiac cells, sensitive to light, and hence capable of polarization and depolarization by the inventive method. At low frequencies of pulse stimulus the cell becomes stronger. However at higher pulse frequencies, i.e., >30 Hz repeated depolarization, or prolongation of the depolarization state, leads to increased vulnerability of these cells in the presence of other toxic agents or insults, such as both medication and increased temperatures applied during hyperthermal therapy. The inventive method thus can effectively "paralyze" the cancer cell and kill it, similar to how a state of prolonged depolarization of brain cells in epilepsy or post-traumatic epilepsy leads to neuronal degeneration and dementia in these patients. This effect can be recreated in cancer cells, either directly by use of a direct electrical pulse, or indirectly by use of light/electromagnetic radiation or ultrasound pulses when cancer cells are in contact with previously administered coated nanoparticles such as quantum dots or piezoelectric nanoparticles or piezoelectric quantum dots.

The ultrasound-created electrical pulse on the nanoparticles is transmitted to adjacent cells, depolarizing the cellular membrane. Simultaneously, the electromagnetic radiation or

light produces an electrical field affecting the cell membrane ion channels or rhodopsin channel, leading to cell membrane depolarization. Interestingly, a biphasic stimulus produced by light or electricity at low frequencies, i.e., <20 Hz to 30 Hz, can be used beneficially in vitro or in vivo to enhance cellular survival or to cause normal cell multiplication in presence of growth factors. The inventive method thus has capabilities to both target destruction of cancer cells, while also to facilitate or augment normal cell capabilities under two different conditions. For stimulation of normal cells, a low frequency pulse is used to render normal cells more viable. For destruction of tumor cells, a high frequency pulse maintains the cell in a state of depolarization, rendering the tumor cell "leaky" and thus more vulnerable to additional external insult or medication.

As an example, in cultured stem cells, a biphasic stimulus causes the stem cells to grow faster and multiply (data not shown). It will be appreciated by those skilled in the art that it is important to differentiate these two processes properly, in order to achieve the desired effect in each situation.

Tumor suppressor genes include, but are not limited to, those encoding p53, APC gene, DPC-4/Smad4 gene, BRCA-1 gene, BRCA-2 gene, WT-1 gene, retinoblastoma gene (Lee et al., Nature, 329: 642 (1987)), MMAC-1 gene, adenomatouspolyposis coil protein, deleted in colorectal cancer (DCC) gene, MMSC-2 gene, NF-1 gene, nasopharyngeal carcinoma tumor suppressor gene that maps at chromosome 3p21.3, MTS1 gene, CDK4 gene, NF-1 gene, NF-2 gene, Rb, Rap1A, DCC, k-rev, and VHL gene.

Therapeutic genes include, but are not limited to, genes 30 encoding enzymes, blood derivatives, hormones, lymphokines such as interleukins, interferons, tumor necrosis factor, etc., growth factors, neurotransmitters or their precursors or synthetic enzymes, trophic factors such as BDNF, CNTF, NGF, IGF, GMF, αFGF, βFGF, NT3, NTS, HARP/pleiotro- 35 phin, etc., apolipoproteins such as ApoAI, ApoAIV, ApoE, etc., dystrophin or a minidystrophin, the CFTR protein associated with cystic fibrosis, intrabodies, coagulation factors such as factors VII, VIII, IX, genes participating in DNA repair, suicide genes defined as genes whose products cause 40 cell death, e.g., thymidine kinase (HS-TK), cytosine deaminase, etc., pro-apoptic genes, prodrug converting genes defined as genes encoding enzymes that convert prodrugs to drugs, a gene encoding MY07A providing therapy for Usher's syndrome (1B), genes encoding somatostatin receptor 45 subtype 2 (sst2) and deoxycitidine kinase uridylmonophosphate kinase (dck::umk), which exhibit complementary therapeutic effects for therapy in advanced and/or metastatic pancreatic cancer due to both genes inducing an antitumor bystander effect and rendering gemcitabine treatment more 50 efficient, and anti-angiogenic genes or alternatively, genes such as VEGF that promote angiogenesis.

Defective genes include, but are not limited to, BACE-1 for therapy of Alzheimer's disease,  $\alpha$ -synuclein for therapy of Parkinson's disease, htt for therapy in Huntingdon's disease, neuronal caspase-3 for therapy of stroke to reduce ischemic damage, etc.

Genes encoding products for synthesis or inhibition of a therapeutic protein include genes encoding anti-cancer agents, growth factors, hypoglycemic agents, anti-angio- 60 genic agents, bacterial antigens, viral antigens, tumor anti-gens, and/or metabolic enzymes. Examples of anti-cancer agents include, but are not limited to, interleukin-2, interleukin-4, interleukin-7, interleukin-12, interleukin-15, interferon- $\alpha$ , interferon- $\beta$ , interferon- $\gamma$ , colony stimulating factor, granulocyte-macrophage stimulating factor, anti-angiogenic agents, tumor suppressor genes, thymidine kinase, eNOS,

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iNOS, p53, p16, TNF-α, Fas-ligand, mutated oncogenes, tumor antigens, viral antigens, and/or bacterial antigens.

One skilled in the art will appreciate that combinations of genes may be used, e.g., genes responding to light to produce action potential, and genes modifying defective genes a tissue or organ to restore normal function of that organ. For example, therapy can be targeted to eye, brain, lung, spinal cord, peripheral nerves, digestive tract, etc., to restore swallowing, breathing, glandular function, etc.

In one embodiment, the gene induces expression of an enzyme with antioxidant activity. In one embodiment, the gene encodes cystic fibrosis transmembrane conductance regulator (CFTR) for therapy of cystic fibrosis. In one embodiment, the gene encodes Bruton's tyrosine kinase for therapy of an X-linked agammaglobulinemia. In one embodiment, the gene encodes an adenosine deaminase (ADA) for therapy of severe combined immunodeficiency (SCID). In one embodiment, the gene encodes Factor IX for therapy of 20 hemophilia B. In one embodiment, the gene encodes SMN-1 for therapy of spinal muscular atrophy. In one embodiment, the gene encodes an anti-VEGF protein (e.g., sFlt-1) for therapy of exudative age related macular degeneration (AMD). In one embodiment, the gene encodes Rab-Escort Protein (REP-1) for therapy for choriodemia. In one embodiment, the gene encodes RPE65, an RPE-specific 65-kDA protein involved in conversion of all-trans retinol to 11-cis retinal during phototransduction, for therapy of the genetic defect in Leber's congenital amaurosis. In one embodiment, the gene encodes MERTK for therapy of retinitis pigmentosa. In one embodiment, the gene encodes ABC4, a a protein involved in energy transport to and from photoreceptor cells in the retina, for therapy of Stargardt's syndrome.

In embodiments, genes are provided for therapy in ocular pathological conditions, some of which were previously described. Examples include, but are not limited to, retinitis pigmentosa, color blindness, wet and dry ARMD, diabetic retinopathy, corneal dystrophies, Meesman syndrome, Fuchs syndrome, granular and macular corneal dystrophies, keratoconous, Sejorgen's syndrome, inherited glaucoma, retinohyaloidopathies, congenital cataract, Marfan syndrome, choridermia x-linked retinoschisis, achromatopsia, etc.

One embodiment of the inventive method is administering coated nanoparticles to a patient to verify that the nanoparticles are in contact with the appropriate tissue, including circulating or non-circulating cells, for photoacoustic technology imaging at relatively low temperatures of 37° C. to 43° C. The image represents the structure that is being treated, e.g., a tumor mass such as cancer, or normal organ tissue such as brain, retina, spinal cord, heart, or any other organ.

In use, a photoacoustic unit controls the thermal energy unit to achieve a precise temperature of 37° C. to 43° C. to release the gene or a drug attached to a thermosensitive polymer. The photoacoustic system controls the thermal energy unit using software that responds to a predetermined program. The program initially provides a low energy level to produce a temperature of 37° C.-42° C. for imaging and drug release that is achieved in less than 30 seconds upon application. The program then provides a high energy level to result in a defined temperature of 44° C.-50° C. to 70° C. or more at the site of the nanoparticle for a very short duration of one femtosecond to less than one second. This high energy melts or opens a hole in the cellular membrane through which the medicament(s) and/or gene(s) that were previously released from the nanoparticles enter the cells to exert an effect. In some embodiments the medicament(s) and/or gene(s) may still be attached to the nanoparticles as it enters the cell.

In the gene therapy embodiment, the desired gene or genes are attached to the nanoparticles. They may be administered using enolase and/or clustered regularly interspersed palindromic repeats (CRISPR), both of which are known in the art. Cas9, the CRISPR associated protein 9, is an RNA-guided DNA endonuclease associated with the CRISPR complex. The endonuclease and/or the CRISPR/Cas9 complex, upon cell entry, seek DNA containing a target sequence, e.g., a specific mutation in the mitochondria or the nucleus. The cleavage functions of the endonuclease and/or CRISRP/Cas9 complex destroy the disease-producing DNA by excising it or replacing it with an un-mutated, i.e., normal DNA segment, as known in the art to treat the patient. The nuclease cleaves host DNA at the target site. With one cut, CRISPR can insert a new customized gene sequence; with two cuts, two CRISPRs can excise the DNA, providing the host with a new DNA. The mechanism of action of the CRISPR/cas9 complex and its ability to target and edit genes is known in the art (e.g., Sander and Joung, CRISPR-Cas systems for editing, regulating and 20 targeting genomes, Nature Biotechnology 32 (2014) 347-355 which is expressly incorporated by reference herein). The sudden temperature rise created with the previously described thermal unit opens the cell membrane causing the genes to enter the cells along with medication. Nanoparticles 25 can be functionalized to render or enhance biocompatibility, and can be further treated or delivered to enhance cell penetration. The nanoparticles can comprise an enolase or CRISPR complex, and/or a gene, and an antibody that targets the nanoparticles to a cell, and can be coated with a biocompatible molecule for cell uptake, forming a complex that is activated with an energy source. As is known in the art, an antibody may be used to target the nanoparticle to a particular cell, cell type, organ, location in the body, etc.

Once inside the cell, the genes along with the CRISPR 35 complex enters the cell nucleus or mitochondria to precisely modify en masse the cellular genetic information, i.e., the gene pool, of a tumor or organ. This either provides therapy for a disease caused by a genetic deficiency, or adds a new useful gene or genes to remove and replace the defective gene 40 or genes in the cell nucleus or mitochondria.

The nanoparticles or quantum dots absorb the energy selectively from an external source and heat up faster than the surrounding structure. The particle material may be ceramic, plastic, silicon; particles of crystalline silicon may be monoc- 45 rystalline cells, poly or multicrystalline cells, ribbon silicon having a multicrystalline structure, nanocrystals of synthetic silicon, gallium/arsenide, cadmium/selenium, copper/indium/gallium/selenide, zinc sulfide, indium/gallium/phosphide, gallium arsenide, indium/gallium nitride, a nanocrys- 50 tal, such as cadmium/selenium (Cd/Se) and a metal, e.g., a CdSe/Au nanometer-sized composite particle as previously described, particles of a variety of semiconductor/metal and semiconductor/semiconductor hetero-junctions, e.g., particles based upon semiconductor/metal hetero-junctions 55 between group II-VI, IV, III-V, IV-VI, referring to groups of the periodic table, metal-oxide, or organic semiconductors and a metal, and in particular those based upon Si/Au, GaAs/ Au, InAs/Au, and PbS/Au hetero-junctions. The quantum dots and/or semiconductor nanowires may also be biocom- 60 patible short peptides of naturally occurring amino acids that have the optical and electronic properties of semiconductor nano-crystals, e.g., short peptides of phenylalanine. The particles can consist of both inorganic or organic materials, as previously described. Nanoparticles, as used herein, include 65 quantum dots, non-quantum dot nanoparticles, and the following structures.

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Graphene quantum dots, graphene-oxide quantum dots, graphene-zinc oxide quantum dots, graphene-zinc oxide quantum nanowires, carbon nanotubes, nanocrystals such as cadmium/selenium (Cd/Se), and particular types of each, e.g., graphene quantum dots, graphene-oxide quantum dots, graphene-zinc oxide quantum dots, graphene nanotubes, and/or carbon nanotubes. These may be non-magnetic, magnetic, paramagnetic, diamagnetic, supramagnetic, non-magnetic, mesoporous carbide-derived carbon, iron oxide nanoparticles with gold, graphene oxide, mesoporous silicon nanostructures, nanoshells, nanorods, nanotubes, nanowires, etc.

Other carbon-based skeletal-type nanostructures including but not limited to fullerenes, bucky balls, micelles and micelle-like structures such as micellar nanoparticles (MNP), liposomes, dendrimers, aptamers, and single-stranded deoxyribonucleic acid- or ribonucleic acid-oligonucleotide aptamer-conjugated or modified nanoparticles. Specific types include, e.g., dendrimers including poly(amidoamine) (PAMAM), poly(amidoamine-organosilicon) (PAMAMOS), poly(propyleneimine) (PPIO), tecto, multilingual, chiral, hybrid, amphiphilic, micellar, multiple antipen peptide, and Frechet-type dendrimers. Nanoparticles may be synthetic, organic (e.g., liposomes), non-organic, non-magnetic, magnetic, paramagnetic, diamagnetic, supramagnetic, non-magnetic, mesoporous carbide-derived carbon, iron oxide nanoparticles with gold, graphene oxide and mesoporous silicon nanostructures, carbon, quantum dots, nanoshells, nanorods, nanotubes, nanowires, quantum dots, etc. Illustrative and non-limiting specific examples also include liposomal nanoparticles, liposome-PEG nanoparticles, micellar polymeric platform nanoparticles, L-adenine nanoparticles, L-lysine nanoparticles, PEG-deaminase nanoparticles, polycyclodextrin nanoparticles, polyglutamate nanoparticles, calcium phosphate nanoparticles, antibody-enzyme conjugated nanoparticles, polymeric lipid hybrid nanoparticles, nanoparticles containing a combination of two-three elements such as gold, gold-iron oxide, iron-zinc oxide, metallic nanoparticles, polylacticglycolic acid nanoparticles, ceramic nanoparticles, silica nanoparticles, silica crosslinked block polymer micelles, albumin-based nanoparticles, albumin-PEG nanoparticles, dendrimer attached magnetic or non-magnetic nanoparticles, etc.

Hybrid quantum dots include but are not limited to graphene/zinc oxide (ZnO) and reduced graphene oxide, or plasmonic nanoparticles coated with reduced graphene oxide, dextran-reduced graphene oxide, etc. Exemplary nonlimiting examples include liposome-PEG nanoparticles, micellar polymeric platform nanoparticles, L-adenine nanoparticles, L-lysine nanoparticles, PEG-deaminase nanoparticles, polycyclodextrin nanoparticles, polyglutamate nanoparticles, calcium phosphate nanoparticles, antibody-enzyme conjugated nanoparticles, polymeric lipid hybrid nanoparticles, nanoparticles containing a combination of two-three elements such as gold, gold-iron oxide, iron-zinc oxide, metallic nanoparticles, polylacticglycolic acid nanoparticles, ceramic nanoparticles, silica nanoparticles, silica crosslinked block polymer micelles, albumin-based nanoparticles, albumin-PEG nanoparticles, dendrimer attached magnetic or non-magnetic nanoparticles, etc.

ZnO added to graphene quantum dots or to a combination of graphene particles and/or carbon nanotubes with a ZnO nanowire or nanorod using an electron gun. In embodiments, particularly those using light to stimulate the described particle, ZnO is useful because it prevents light reflecting off the particle surface, i.e., it serves as an anti-reflective coating, and provides a more efficient quantum dot compared with graphene or a carbon nanotube alone and ZnO additionally

has the benefit of being an antibacterial compound and thus can be utilized for transporting biomolecules, such as DNA, along with other polymers. These may contribute a further therapeutic function and/or to biocompatibility. Nanoparticles may be coated with biocompatible mono- or bilayers of 5 phospholipid a protein, a peptide polyethylene glycol (PEG) that can be used as a scaffold to aid in biocompatibility of the particle. The particles can be entirely or partially biodegradable. Nanoparticles may also be included in/coated on a bioabsorbable, or non-bioabsorbable but biocompatible, polymer fiber, tube, or other two- or three-dimensional structure. Therapeutic stimulation of the polymer and adjacent tissue may stimulate and/or inhibit cell excitation, depending upon the wavelength of the applied light and the character of the associated particles, with differing parts of the polymer, e.g., 15 the front and back sides of a substantially two-dimensional structure, having different particles to exert different effects upon target and adjacent cells. Optional therapeutic or biocompatibility enhancing molecules include peptides, biotinstreptavadin, (poly)ethylene glycol (PEG), acetyl cysteine, 20 cell penetrating peptide (CPP), arginine-CPP, cysteine-CPP, etc. Nanoparticles can be functionalized to render or enhance biocompatibility, and can be further treated or delivered to enhance cell penetration. The nanoparticles can comprise an enolase or CRISPR complex, and/or a gene, and an antibody 25 that targets the nanoparticles to a cell, and can be coated with a biocompatible molecule for cell uptake, forming a complex that is activated with an energy source.

Nanoparticles can be administered systemically, or applied locally, e.g., as a drop, injected locally, or applied on or with 30 a biocompatible substrate at a site. Therapeutic molecules such as drugs and/or biologics may be contained in or on the nanoparticles.

Nanoparticles may be included in or coated on a bioabsorbable or non-bioabsorbable but biocompatible polymer 35 structured or configured as a fiber, a tube, a substantially two-dimensional structure, or a three-dimensional structure to fit any anatomical or physiological site, configured as any desirable length or size to maintain its position with respect to an anatomical and/or physiological location, e.g., the eye. 40 Upon activation, cells and adjacent tissue may undergo excitation, which is further tunable, e.g., parts of the polymer, e.g., the front and back sides of a substantially two-dimensional structure, having different particles to effect target cells, adjacent cells, etc.

Nanoplatforms may be developed of such carriers with both superior biocompatibility and superior penetrability. Pan et al., ACS Appl. Mater. Interfaces 5 (2013) 7042, disclose one such example of a family of silica crosslinked micelles that are biocompatible, luminescent, and stable; specifically, silica crosslinked pluronic F127 (difunctional block copolymer surfactant) micelles loaded with decyl capped silicon nanoparticles. Erogbogbo et al. (Integr. Biol. 5 (2013) 144) disclose another such example of a biocompatible silicon quantum dot F127 micelle with multiple silicon quantum dots incorporated into a micelle core having gold deposited on the nanostructure surface to provide light scattering properties facilitating imaging, while the silicon nanocrystals maintain photoluminescence.

Nanoparticles may have enhanced cell penetration by 60 being coated on their surface with, encapsulated by, or otherwise associated with polymers have enhanced cell penetrating ability. Synthesis, fabrication, modifications, are possible, e.g., Tomczak et al., Progress in Polymer Science, 34 (2009) 393. As one example, PEG grafted polyethylenimines 65 (PEI) encapsulate and solubilize luminescent quantum dots through direct ligand-exchange reactions via positive charges

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and a proton sponge effect. PEG improves nanoparticle stability and biocompatibility, reduces PEI toxicity to cells, and facilitates cell penetration. Duan and Nle, J. Am. Chem. Soc. 129 (2007) 3333. As one example, nanoparticles can be coated with proton-absorbing polymers, termed proton sponges, to facilitate siRNA delivery by enhanced cell penetration as well as other means. Yezhelyev et al., J. Am. Chem. Soc. 130 (2008) 9006. As one example, the linear polysaccharide chitosan, in the form of beads, gels, sponges, membranes, scaffolds, etc. may be used as PEG-chitosan and/or folate-chitosan quantum dots. Rajan and Raj, I.Re.CH.E. 5 (2013) 145. As one example, folate conjugated, PEG-coated quantum dots specifically recognized and were internalized by folate receptors that were overexpressed in certain cancer cells, i.e., specific receptor-mediated cellular internalization. Song et al., Clincal Chemistry 55 (2009) 955. As one example, PEG-conjugated chitosan derivatives were synthesized and demonstrated narrow size distribution, good water solubility, low cytotoxicity. Lv et al., Chemical Papers 67 (2013) 1404. Various coatings and encapsulation conditions, components, linkage types, etc., i.e., nanoparticle architecture, can assist in regulating nanoparticle penetration effects and parameters under various conditions. Smith et al., Advanced Drug Delivery Reviews 60 (2008) 1226. As one example, at least two of the same or different, i.e., hybrid, nanoparticles can be encapsulated in amphiphillic block copolymer micelles that are crosslinked and thus form a shell surrounding the plurality of nanoparticles. Kim and Taton, Langmuir, 23 (2007) 2196. As one example, nanoparticles such as quantum dots can be encapsulated in carboxylated Pluronic (i.e., poly(ethylene oxide) (PEO) and poly(propylene oxide) (PPO) with a PEO-PPO-PEO structure) 127 triblock, forming a polymeric micelle with efficient cell penetration. Liu et al. Theranostics 2 (2012) 705.

Nanoparticles may be conjugated with physiologic biomolecules to enhance or facilitate cell penetration. Examples of such compounds include, but are not limited to, carbohydrates, cholesterol, glutathione (e.g., taking advantage of amino functionalities), collagen, chitosan, alginate, fibrin, fibrinogen, cellulose, starch, collagen, dextran, dextrin, fragments and derivatives of hyaluronic acid, heparin, fragments and derivatives of heparin, glycosamino glycan (GAG), GAG derivatives, polysaccharide, elastin, and biotin, proving good stability and cell viability, e.g., Jin et al., Int. J. Mol. Sci. 9 (2008) 2044. One example of enhanced penetration is quantum dots conjugated or otherwise associated with cell penetrating agents such as cell penetrating peptides (CPP) and activatable-cell penetrating peptides (ACPP). CPP and ACPP labeled with fluorescent polycationic CPP coupled by a cleavable linker to a neutralizing peptide have been used to visualize tumors during surgery. ACPP conjugated to dendrimers (ACPPDs) and gadolinium chelates can allow MRI visualization of whole body tumors, permitting therapy if the magnetic or gadolinium nanoparticles are labeled with ACPPD. A nanoparticle may be cleavably conjugated to an ACPP by, e.g., a linker such as an ethylene glycol or PEG moiety. It may be contain a label such as a fluorescent moiety, chemiluminescent moiety, etc. The ACPP may be labeled with a polycationic CPP. ACPP and CPP may be naturallyoccurring or artificially constructed protein segments (<30 amino acids) rich in arginine, lysine, cysteine, histidine, ornithine, etc.; preferably  $\alpha$ -helices and about 17-amino acids. The ACPP and CPP may include a penetration accelerating peptide sequence (Pas) or an INF7 fusion peptide sequence. CPP and/or ACCP can be linked to cargoes either covalently or noncovalently, or can use block copolymers to form various kinds of micelles. Liu et al., Endocytic Trafficking of

Nanoparticles Delivered by Cell-penetrating peptides Comprised of Nona-arginine and a Penetration Accelerating Sequence, PLOS ONE 8 (2013) e67100; Liu et al., Intracellular Delivery of Nanoparticles and DNAs by IR9 Cell-penetrating Peptides, PLOS ONE 8 (2013) e6405; Liu et al., Cell-Penetrating Peptide-Functionalized quantum Dots for Intracellular Delivery, J. Nanosci Nanotechnol 10 (2010) 7897; Liu et al., Cellular Internalization of Quantum Dots Noncovalently Conjugated with Arginine-Rich Cell-Penetrating Peptides, J. Nanosci Nanotechnol 10 (2010) 6534; Xu et al., Nona-Arginine Facilitates Delivery of Quantum Dots into Cells via Multiple Pathways J. Biomedicine and Biotechnology volume 2010, Article ID 948543, 1-11. Exemplary but non-limiting ACPP and CPP include transportan, penetratin, TAT, VP22, MAP, KALA, ppTG20, proline-rich peptides, MPG-derived peptides, Pep-1, nona-arginine, and the carboxy-terminal tail of TFPI-2, polyproline helices having cationic amino acids and/or cationic-functionalized amino acids within the helix (e.g., guanidinium-functional- 20 ized proline), a synthesized oligoarginine cell penetrating peptide (based on the HIV-1 Tat protein motif) bearing a terminal polyhistidine (His8) tract facilitated transmembrane ° (Delehanty et al., Bioconjug Chem 17 (2006) 920). Other cell penetrating agents include oligoguanidinium scaffolds, 25 cationic-functionalized calixarenes, or cyclodextrins, e.g., arginine-functionalized calixarene). For example, CPP comprising a INF7 fusion peptide and nona-arginine produces IR9/QD and IR9/DNA complexes, IR9, IR9/QD and IR9/ DNA. Handbook of Cell Penetrating Peptides, Langel (Ed), 30 2007 (Second Edition) CRC/Taylor & Francis). Nanoparticles may be coated or otherwise associated with organic or non-organic biodegradable compounds including but not limited to silicon, porous silicon, aliphatic biodegradable polymers, etc.

Agents may be linked to, associated with, complexed or conjugated with nanoparticles using linking agents and methods known in the art including but not limited to the following: amino groups, carboxyl groups, S—S deprotected sulf-hydril groups in biomolecules, e.g., bis(succinimide 40 derivative conjugation, maleimidosuccinimide/succinimidylpyridyldithio/-halosuccinimide derivative conjugation, N-(4-maleimido-phenyl) isocyanate conjugation, carbodiimide conjugation, sulfosuccinimidylsuberyl linkage, synthetic tripyrrole-peptide linkage, NHS-esters and other esters, etc. 45 Cleavage of the linkers by specific proteases, e.g., matrix metalloproteinase-2, dissociates the polyanion and enables arginine-rich CPPs to enter cells.

As previously described, nanoparticles function as carriers and deliver a gene or genes from a variety of gene families. As 50 known in the art, a gene can integrate into the cell nucleus and nucleolus, and can be delivered alone, or in combination with other genes to correct a genetic defect or to provide functional therapy, e.g., induction of an action potential in a membrane of excitable cells such as retinal cells, cardiac cells, and cells 55 of the central nervous system (CNS) or peripheral nervous system (PNS). Other desired gene stimulators, e.g., promotors, or gene silencers, e.g., siRNA, may also be included to up- or down-regulate gene function.

Nanoparticles may be incorporated within liposomes and/ 60 or plasmids carrying DNA, RNA, siRNA, medications, etc.

Nanoparticles may of any shape, e.g., spheres, nanotubes, nanowires, tetragons, hexagons, cylinders, etc.

Nanoparticles may be coated with PEG, PEI, chitosan, biotin, streptavidin, CPP, ACPP etc. along with the specific 65 cell membrane antibodies that can be tested by enzyme linked immunoassay (ELISA) to attach to the cell receptors.

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For gene transfer, a plasmid containing a gene is attached to the nanoparticles during the coating process with PEG, chitosan, etc. along with CPP or ACPP. Nanoparticles generally may have improved tolerance in vivo compared to quantum dots, and use of magnetic nanoparticles is desirable for gene transfer because a magnetic field increases the transfection in tissue. Magnetic nanoparticles are administered along with a localized magnet. The electrostatic potential of these nanoparticles can be as high as -25 mV, encouraging further cell membrane penetration. While magnetic nanoparticles may be desirable for some embodiments, non-magnetic nanoparticles and organic nanoparticles can also be used with plasmids for transfection. In one embodiment using nanoparticles for gene transfer, a liposome containing a gene forms a nanoparticle-liposome-gene complex. Liposomes as a carrier for the nanoparticle and gene are then coated and conjugated with antibody prior to administration to the patient. Methods of preparing liposomes are known in the art, e.g., Akbarzadeh at al., Nanoscale Res. Lett. 8 (2013) 102, which is expressly incorporated herein by reference in its entirety.

Piezoelectric nanoparticles may be stimulated by a wireless, remotely controlled device such as an ultrasound probe to produce an electrical current that may result in altered membrane potential, i.e., hyperpolarization, hypopolarization, or depolarization, or an altered action potential of cells in contact with the piezoelectric nanoparticles. The piezoelectric nanoparticles may be administered by injecting submucosally and/or spraying a nasal mucosa to travel through olfactory nerves to the brain, and the brain stimulated with an externally applied ultrasound over the nose by a patient as needed. Piezoelectric nanoparticles can be rendered biocompatible by coating with zirconate titanate, perovskite-based, oxides, barium titanate, polyvinylidene fluoride (PVDF), etc. The piezoelectric nanoparticles may be controlled by a diode laser surgically implanted in the patient, with the piezoelectric nanoparticles acting as an "on"/"off" switch of a battery supplying electricity to the diode laser. Stimulation is controlled by a programmable processor of an extra-corporal ultrasonic probe wirelessly controlled by the diode laser, which is implanted, e.g., external to the skull, internal to the skull, adjacent to an organ, adjacent to sympathetic ganglion cells, adjacent to parasympathetic ganglions cells, adjacent to the brain, adjacent to the heart, adjacent to an eye, adjacent to the spinal cord, adjacent to peripheral nerves, etc. In this example, the patient can be treated for epilepsy, migraine, depression, paresis, paralysis, spastic muscles numbness, nerve injuries, dementia, spinal cord injury, peripheral nerve injury, Parkinson's disease, Alzheimer's disease, restless leg syndrome, etc. A ribbon of piezoelectric nanoparticles may be attached to the battery of the diode that, upon ultrasound stimulation, creates a pulsating electrical current in the piezoelectric nanoparticles leading to the battery of the diode laser, thus controlling activation of the diode laser light production and initiating an "on"/"off" action potential in the membrane of excitable cells depending on the frequency and duration of the electrical pulse.

Nanoparticles can be coated such that the nanoparticle is amphiphilic, where a portion of the nanoparticle is rendered hydrophilic and another portion of the nanoparticle is rendered hydrophobic. For example using a CdSe/Au particle, the CdSe/Au particles are covered by trioctylphosphine oxide and alkylphosphonic acid, both of which are hydrophobic. Surface functionalization covers the Au portion of the CdSe/Au particles with PEG, making them hydrophilic; the CdSe portion, still covered by trioctylphosphine oxide and alkyl phosphonic acid, remains hydrophobic. CdSe/Au particles may be suspended in N,N-dimethylformamide containing

detergent (e.g., 1% Triton X-100) and exposed to polyethylene glycol-( $\mathrm{CH_2}$ ) $_{10}$ —SH to coordinate the thiol to the Au end. The nanoparticles may be coated with or linked to, e.g., folate, polydopamine, etc. so that these molecules are targeted intracellularly, extracellularly, to a cell membrane, to a 5 specific cellular site or organelle, etc. The nanoparticles may be coated with a thermosensitive polymer, e.g., chitosan, PEG, etc. Application of an external energy source results in a slight increase in temperature, e.g., to about 39° C. to about 43° C. in one embodiment, to about 40° C. to about 42° C. in 10 another embodiment, that facilitates release of the biomolecule from the thermosensitive nanoparticles. The nanoparticles can be made to respond to various wavelengths of light (visible and invisible). In one embodiment they are coated with organic molecules. In one embodiment, they are com- 15 pletely organic. In one embodiment, they are PEGylated to last longer. In one embodiment, they are coated to be attracted to certain receptors or stay only on the cell surface.

In one embodiment one administers systemically, in a body cavity, or locally, a plurality of nanoparticles, which can 20 include quantum dots, or plasmonic nanoparticles. The nanoparticles may be magnetic, paramagnetic, nonmagnetic, metallic, non-metallic, organic, inorganic, synthetic, piezoelectric, nanobots, lanthanide, cerium, gold, zinc, silver, silicon, perovskite; or liposomes, dendrimers, nanotubes, 25 nanowires, caged nanoparticles, nanoshells, etc. The nanoparticles may be coated with a biocompatible polymer such as biotin, streptavidin, PEG, cell penetration peptide (CPP), arginine CPP, etc., known in the art. The nanoparticles may be conjugated with antibodies or aptamers for cell targeting.

The size of the nanoparticles ranges from 1 nm to greater than 999 nm. A nanoparticle size in the range of 1 nm to 20 nm is ideal for the eye and CNS to access the intercellular space and travel to and clear the kidney without resulting in systemic side effects. The location of the organ prescribes the 35 source of energy used that can access external or internal structures. One can use an external light source for, e.g., the eye, skin, mucosa, intestinal tract, airways, etc. Internal organs can be reached with a mechanical source of energy such as ultrasound, etc. For internal organ interiors, one can 40 use a fiber optic, or one may select nanoparticles that can be stimulated by another source of energy that easily travels through tissue, e.g., radiofrequency waves, microwaves, alternating magnet, and ultrasound. Energy sources includes, but are not limited to, electromagnetic radiation, light both 45 visible and invisible, alternating magnet, reversible magnet,

The nanoparticles expand after exposure to the source of energy, creating a photoacoustic sound wave that can be recorded by an ultrasound receiver, amplified, converted to an 50 electrical signal, and imaged in one, two, or three dimensions. Any of the following imaging modalities may be used, either alone or in combination: photoacoustic imaging, nanoparticle imaging, super resolution imaging, anatomical imaging, vascular imaging, functional imaging, molecular imaging, tumor 55 imaging, photoacoustic computed tomography, ultrasound, focused ultrasound, magnetic resonance imaging (MRI), functional MRI (fMRI), computed tomography (CT), positron emission tomography (PET), OCT, and alternating magnetic field imaging, resulting in enhanced image acquisition, resolution, visibility, distinction, and/or utility.

The nanoparticles may be administered by any route, including but not limited to injection into tissues, organs, or body cavities, systemic injection, intravenous injection, local injection, submucosal injection, mucosal spraying (e.g., nasal 65 mucosa for travel through the olfactory nerves to various areas of the brain that can be stimulated with an externally

applied ultrasound over the nose by a patient as needed e.g. in epilepsy, migraine, depression, Parkinson's disease, Alzheimer's disease, restless leg syndrome, etc.), oral administration, topical administration to skin and mucous membranes, etc. In one embodiment, the nanoparticles are injected into the circulatory system. While other administration methods are within the inventive scope, the ease and convenience of direct intravenous injection, or less routinely intraarterial, injection makes this route attractive and provides a rapid route to reach all organs. Topical, intrathecal, subcutaneous, submucosal, inhalation, oral, cerebrospinal fluid injection, or intracavity (bladder, uterus, etc.) administration routes are possible to directly reach tumors or lymph nodes.

The various aspects of the invention are basically attained by providing a method of hyperthermally treating tissue in an animal. The method comprises the step of introducing a temperature indicating substance into the bloodstream of the animal to flow through a target site. The temperature indicating substance includes a fluorescent dye encapsulated within a heat sensitive liposome. The fluorescent dye is releasable from the liposome at a temperature of at least 41° C. A heat source is applied to the target site and the target is hyperthermally heated to at least 41° C. to release and fluoresce the dye and to hyperthermally treat the target site for a time sufficient to kill cells in the tissue.

The aspects of the invention are also attained by providing a method of detecting a threshold temperature and of hyperthermally treating tissue in an animal. The method comprises the step of introducing a first fluorescent dye encapsulated in a first heat sensitive liposome into the bloodstream of an animal in a location to flow through a target site in the animal. The first fluorescent dye is releasable from the first heat sensitive liposome at a temperature of at least 41° C. The target site is heated to a temperature to release the first fluorescent dye and the first fluorescent dye is fluoresced to indicate and visualize a tissue temperature of at least 41° C. Heating of the target site is continued at a temperature of at least 41° C. for a time sufficient to hyperthermally treat the tissue.

Another aspect of the invention is a method of targeting tumor stem cells for therapy in a patient needing the therapy, e.g., a cancer patient either simultaneously undergoing chemotherapy or having previously undergone chemotherapy. The method administers chemotherapy with thermotherapy, with thermotherapy being a stepwise increase in temperature that, at relatively lower temperatures, primes the tumor cells, including tumor stem cells, for increased susceptibility to the chemotherapy. Then at relatively higher temperatures, the method kills the primed tumor stem cells with a synergistic combination of the chemotherapy and the thermotherapy.

Another aspect of the invention is a method of targeting tumor cells, tumor stem cells, and endothelial cells of tumor-associated vessels to selectively destroy both the tumor as well as the vessels feeding the tumor by combining thermal therapy with localized internally administered ionizing radiation therapy.

The aspects of the invention are further attained by providing a method of hyperthermally treating tissue of an animal. The method comprises the step of introducing a temperature indicating substance into the bloodstream of the animal to flow through a target site. The temperature indicating substance includes a first fluorescent dye encapsulated in a first temperature sensitive liposome. The first fluorescent dye is releasable from the first liposome by heating to a temperature of at least 42° C. A second fluorescent dye encapsulated in a second temperature sensitive liposome is also included. The second fluorescent dye is releasable from the second lipo-

some by heating to a temperature of at least  $50^{\circ}$  C. The target site is heated to a temperature of at least  $42^{\circ}$  C. The first fluorescent dye is fluoresced to indicate an effective temperature for hyperthermally treating the tissue without releasing the second fluorescent dye from the second liposomes.

Another aspect discloses a method of delivering particulate material, e.g., nanoparticles, into mammalian cells and/or tumor cells through pinocytic uptake and/or using various channels. In one embodiment, the particulate material is delivered to the nuclear membrane, which typically has a pore 10 size <100 nm in diameter. In one embodiment, the pores and/or pinocytic vesicles etc. permit the nanoparticles, smaller than 100 nm, to be taken up into the cytoplasm or the nucleus of the cells/bacteria. The nanoparticles are coated with or contain an antibody and/or drug. The non-magnetic 15 and/or magnetic nanoparticles, once taken up by the cell/bacteria, are heated selectively by an external source such as electromagnetic energy or a reversible magnetic field, and imaged using either a photoacoustic technology or MRI.

One embodiment is a method of providing therapy by 20 administering, to a patient in need of therapy, nanoparticles coated or otherwise associated with an antibody to specific cells, under conditions sufficient to permit antibody accumulation at a tissue target site, radiating the target site with an energy source to penetrate into the tissue target site to con- 25 trollably heat the nanoparticles and generate thermal energy to induce a photoacoustic signal or sound wave from the nanoparticles, using a processor to control the amount of thermal energy delivered at the desired temperature to the target site, recording the temperature and photoacoustic signal or sound wave from the target site or from one or more multiple locations, and amplifying and processing the recorded photoacoustic signal or sound waves to generate a computational tomographic image of the nanoparticles at the tissue target site.

In this embodiment, imaging generates high-resolution two- or three-dimensional photoacoustic images. In one embodiment, the antibody is an anti-tumor antibody, the nanoparticles are attached to tumor cells, and the method generates a two- or three-dimensional photoacoustic image of 40 the tumor. In one embodiment, the antibody is directed to a receptor on a normal cell of an organ, the nanoparticles are attached to the normal cells, and the method generates a twoor three-dimensional photoacoustic image of an organ regardless of location of the organ in the body. Imaging may use 45 radiofrequency, microwave, ultrasound, and/or focused ultrasound. Imaging may be by photoacoustic temperature imaging combined with ultrasound, focused ultrasound, magnetic resonance imaging (MRI), functional MRI (fMRI), computed tomography (CT), positron emission tomography (PET), 50 OCT, and alternating magnetic field imaging, resulting in enhanced image acquisition, resolution, visibility, distinction, and/or utility.

In this embodiment, the processor communicates the temperature from the photoacoustic source to the energy source 55 to control the amount and duration of energy delivered to a desired temperature. The photoacoustic signal or sound wave is recorded and produced as each of a thermal graph of the target site, and as an image in one-dimension, two-dimensions, or three-dimensions.

In this embodiment, the energy source to radiate the target site may be electromagnetic radiation, ultraviolet radiation, visible light, infrared light, radiofrequency waves, microwaves, focused ultrasound, and/or alternating magnetic field radiation. A combination of energy sources decreases the 65 amount of energy required for each unit or source, e.g., lasers of various wavelengths, radiofrequency waves, microwaves

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with focused ultrasound, and microwaves with alternating magnets. Energy is applied from multiple sites or using multiple energy sources thus minimizing or preventing pain to the patient and overheating of the target site. The applied energy may be continuous, intermittent, oscillatory, and/or pulsed. Energy applied in an oscillatory or pulsed manner reduces thermal damage to normal cells while sufficiently heating cells to which the nanoparticles are attached to the desired temperature. Energy may be applied intermittently as an alternating magnetic force to heat to the nanoparticle-cell complex, and imaging measures the temperature of the heated tissue and images the target site. In this embodiment, supramagnetic ferric oxide and/or supraparamagentic ferric oxide nanoparticles, nanotubes, and nanowires are particularly preferred.

In this embodiment, the target site may be maintained at temperatures ranging from  $39^{\circ}$  C. to  $48^{\circ}$  C.,  $37^{\circ}$  C. to  $40^{\circ}$  C.,  $42^{\circ}$  C. to  $46^{\circ}$  C.,  $47^{\circ}$  C. to  $50^{\circ}$  C., and  $50^{\circ}$  C. to  $58^{\circ}$  C.

In one embodiment, nanoparticles are attached to tumor cells, the tumor cells are selectively or preferentially treated over normal non-tumor cells, and the thermal energy is applied from multiple locations covering an area of  $1^{\circ}$  to  $90^{\circ}$ , up to  $180^{\circ}$ , or greater than  $180^{\circ}$  of the circumference of the target tissue.

The patient undergoing therapy by this embodiment of the method may have a benign or malignant lesion, and/or an infection. Cells may be preloaded stem cells, macrophages, dendritic cells, and/or exosomes of dendritic cells, with the method locating or tracking the cells throughout the body, and imaging the cells accumulation in vessels or tissues at a site of inflammation.

In this embodiment the nanoparticles may be synthetic, organic, non-organic, non-magnetic, magnetic, paramag-35 netic, diamagnetic, supramagnetic, non-magnetic, mesoporous carbide-derived carbon, iron oxide nanoparticles with gold, graphene oxide and mesoporous silicon nanostructures, carbon, quantum dots, nanoshells, nanorods, nanotubes, nanobots, and/or nanowires. The nanoparticles may be liposomal nanoparticles, liposome-PEG nanoparticles, micellar polymeric platform nanoparticles, L-adenine nanoparticles, L-lysine nanoparticles, PEG-deaminase nanoparticles, polycyclodextrin nanoparticles, polyglutamate nanoparticles, calcium phosphate nanoparticles, antibody-enzyme conjugated nanoparticles, polymeric lipid hybrid nanoparticles, nanoparticles containing a combination of two-three elements such as gold, gold-iron oxide, iron-zinc oxide, metallic nanoparticles, polylacticglycolic acid nanoparticles, ceramic nanoparticles, silica nanoparticles, silica crosslinked block polymer micelles, albumin-based nanoparticles, albumin-PEG nanoparticles, and/or dendrimers attached magnetic or nonmagnetic nanoparticles. In one specific embodiment, nanoparticles are iron oxide coated with an oligosaccharide. Nanoparticles may be coated with thermosensitive polymers carrying agents, e.g., anti-infectives, chemotherapeutics, anti-VEGFs, anti-EGFRs, hormones, immunosuppressants, immunostimulatory agents, DNA, RNA, siRNA, genes, and/ or nucleotides. The nanoparticles may be coated or otherwise associated with biocompatible molecules, e.g., PEG, biotin, CPP, ACPP, dendrimers, dendrimers conjugated with poly beta amine, and/or small organic molecules. The nanoparticles may be lanthanide, cerium, gold, zinc, silver, silicon etc. The nanoparticles may be perovskite nanoparticles, liposomes, dendrimers, nanotubes, nanowire, caged nanoparticles, etc. All non-biocompatible nanoparticles are coated with a biocompatible polymer such as biotin, streptavidin, (poly)ethylene glycol (PEG), cell penetration peptide (CPP),

arginine CPP, etc., as known in the art, and are then conjugated with antibodies or aptamers for cell targeting.

In one illustrative and non-limiting embodiment, targeted dendrimer nanoparticles or liposomes of, e.g., 20 nm-100 nm size, are conjugated with PEG/gene and an aptamer or an 5 antibody for cell targeting, coated with a heat-sensitive polymer such as chitosan or any others with (list below). These nanoparticles are administered simultaneously with ferric oxide nanoparticles of 1 nm-10 nm size to deliver medication, and are heated with an energy source under the control of a 10 photoacoustic unit to achieve a desired temperature in tumor cells. The temperature is kept initially at 38° C.-43° C. for release of the the gene/medicament, and/or increased to 44° C.-58° C. to damage the cell membranes.

In this embodiment, the antibody may target a circulating cell or a a non-circulating cell fix in an organ or specific location, and the cell may be either benign or malignant. The antibody may be, e.g., abciximab, adalimuab, alemtuzumab, brentuximab, blimubab, canakinumab, alemtuzumab, cituximab, cetrolizumab, daclizumab, denosumab, efalizumab, 20 gemtizumab, glimumab, lipilimumab, infliximab, rituximab, muromonab, oftamumab, palivizumab, panitumab, ranibizumab, tositumomab, and/or rastuzumab.

In a specific embodiment of this general embodiment, the method is provided to the patient with a method for enhanced 25 cell penetration of the antibody-nanoparticle complex, e.g., application of ultrasound and/or electroporation. In another specific embodiment, the method uses a combination of CPP and poly beta amine to enhance cell penetration of the antibody-nanoparticle complex. This embodiment may reduce 30 the total amount of energy applied to achieve therapy of cells at the target site while sparing normal cells.

In another specific embodiment of this general embodiment, the patient is undergoing cancer immunotherapy and the method provides an agent to the patient to result in enhanced immunogenicity, e.g., an inhibitor to CHK1, CHK2, CHM1, and/or CHM2.

43° C.-45° C., greater than 42° C. and up to 58° C. The complex may than 42° C. and up to 58° C. The complex may therefore the patient of this general endough the aptamer and nanoparticle and/or quantum despite the aptamer and appears and a

In another specific embodiment of this general embodiment, patient cells are grown in culture with the nanoparticles, and the cultured cells with nanoparticles incorporated 40 are injected into the patient. The cells grown in culture may be stem cells, macrophages, dendritic cells, lymphocytes and other leukocytes, and/or exosomes from dendritic cells. The nanoparticles can be imaged and traced in the patient, e.g., one or several organs, and can result in indirectly imaging an 45 inflammatory process in tissues of the organ.

The nanoparticles size may range from 1 nm-800 nm, 20 nm to 200 nm, 1 nm-20 nm, or 1 nm-10 nm.

In another specific embodiment of this general embodiment, the target site contains a lesion or pathology, and the 50 nanoparticles contain a polymeric coating that itself contains a medicament that is released when the temperature is 40° C.-47° C. resulting in combined thermal and medicament therapy to the lesion.

The another specific embodiment of this general embodiment, the patient undergoing therapy is treated simultaneously or substantially simultaneously with hemofiltration, hemoadsorption, mesoporous carbide-derived carbon, or another type of agent to prevent a cytokine storm.

In another specific embodiment of this general embodiment, photoacoustic imaging in combination with MRI, ultrasound, or light results in enhanced image distinction of a structure without heating the structure significantly beyond 37° C.-39° C.

In another specific embodiment of this general embodi-65 ment, the method creates a photoacoustic image of an internal structure with radiofrequency wave energy or microwave 22

energy alone. An antibody-coated nanoparticle is targeted to a tumor or structure that, because nanoparticles of 1 nm are significantly smaller than a micron, two nanoparticles are separated only by <2 nm-3 nm, using thermal energy, creating distinct separate signals that are electronically enhanced and, using photoacoustic technology, two points separated by that distance are calculated and measured. This provides enhanced imaging beyond optical resolution of a structure using imaging by, e.g., light, MRI, CT, and/or PET.

One embodiment is a method of providing therapy to a patient requiring therapy by administering a complex of nanoparticles and/or quantum dots that are associated with at least one aptamer that targets the nanoparticles and/or quantum dots to a site under conditions that permit complex accumulation at the target site. An energy source is then provided to the target site to penetrate the tissue and controllably heat the nanoparticles and/or quantum dots and generate thermal energy to induce a photoacoustic signal or sound wave from the nanoparticles and/or quantum dots. The method uses a processor to control the amount of thermal energy delivered at the desired temperature to the target site. The temperature is then recorded with the photoacoustic signal or sound wave from the target site, or from one or more multiple locations, and the recorded photoacoustic signal is amplified and processed to generate a computational tomographic image of the nanoparticles and/or quantum dots at the target site. The energy source may be electromagnetic radiation, ultrasound, radiofrequency waves, microwave energy, focused ultrasound, a magnetic field, a paramagnetic field, an alternating magnetic field, etc. Hyperthermal therapy resulting in cell membrane damage may be achieved a temperature between  $43^{\circ}$  C.- $45^{\circ}$  C., greater than  $42^{\circ}$  C. and up to  $47^{\circ}$  C., or greater than 42° C. and up to 58° C. The complex may be adminis-

In one specific embodiment of this general embodiment, the aptamer and nanoparticle and/or quantum dot complex may contain an agent and a thermosensitive polymer that releases the agent at the target site when a pre-defined temperature at the target site is reached. The agent may be, e.g., a medicament, a biologic, a gene, RNA, RNAi, siRNA, DNA, etc. The specific temperature to release the agent is in the range of 41° C. to 42° C. inclusive. For example, an antivascular endothelial growth factor (anti-VEGF) agent may be administered to a patient with an ocular disease.

In embodiments where the complex contains a gene, an opsin family gene may be administered to an excitable cell having a genetic defect in the opsin family gene and, upon energy stimulation, the opsin family gene administered causes an action potential in the excitable cell membrane. Examples of excitable cells are known to those skilled in the art and include, but are not limited to, retinal cells, brain cells, spinal cord cells, cardiac cells, etc.

In one specific embodiment of this general embodiment, the aptamer and nanoparticle and/or quantum dot complex may contain a photosensitizer to effect phototherapy, in addition to hyperthermal therapy.

In one specific embodiment of this general embodiment, the aptamer and nanoparticle and/or quantum dot complex may contain a radionuclide to effect radiotherapy, in addition to hyperthermal therapy.

In one specific embodiment of this general embodiment, the aptamer and nanoparticle and/or quantum dot complex may contain an immune stimulating agent to effect immunologic therapy, in addition to hyperthermal therapy. The immune stimulating agent may be, e.g., CHK1, CHK2, CHM1, CHM2, etc.

In one specific embodiment of this general embodiment, the aptamer and nanoparticle and/or quantum dot complex may also contain antibodies to target the nanoparticles and/or quantum dots to the tissue target site. For example, anti-beta amyloid and/or anti-Tau protein antibodies may be injected 5 locally into the cerebrospinal fluid or systemically to a patient with Alzheimer's disease.

In one specific embodiment of this general embodiment, imaging generates a high-resolution two- or three-dimensional photoacoustic image that can be optionally overlaid with another image produced simultaneously by, e.g., magnetic resonance imaging (MRI), magnetic resonance spectroscopy (MRS), Raman spectroscopy, ultrasound, focused ultrasound, bioluminescence, optical fluorescence, functional MRI (fMRI), computed tomography (CT), positron semission tomography (PET), OCT, alternating magnetic field imaging, molecular Imaging (MI), imaging using contrast agents, diffusion sensitive magnetic resonance imaging, etc.

In one specific embodiment of this general embodiment, the aptamer and nanoparticle and/or quantum dot complex 20 may be coated or otherwise associated with biocompatible molecules, e.g., (poly)ethylene glycol (PEG), biotin, cell penetrating peptides (CPPs), ACPP, dendrimers, dendrimers conjugated with poly beta amine, small organic molecules, etc. For example, when is complex is to be administered systemically or not locally, the complex may have at least a partial coating of PEG at a thickness sufficient to minimize or prevent damage by plasma enzymes to DNA and/or RNA contained in the complex. Local injection at a site internal to the blood brain barrier results in minimized plasma enzyme degradation of any RNA and/or DNA contained in the complex.

In one specific embodiment of this general embodiment, the method may be used with a method for enhanced cell penetration of the aptamer and nanoparticle and/or quantum dot complex. The method for enhanced cell penetration of the 35 complex may be ultrasound and/or electroporation. This specific embodiment results in therapy of cells at the target site while sparing normal cells.

One general embodiment is a method of enhancing a hyperthermal therapy benefit to a patient in need thereof by 40 selecting at least one nanoparticle among a plurality of nanoparticle types, nanoparticle components, nanoparticle complexes, and nanoparticle compositions; optionally selecting a therapeutic agent and/or biological agent to be carried by the nanoparticles; selecting among a plurality of energy types to 45 activate the nanoparticles by electromagnetic radiation; optionally selecting among additional agents to perform a function separate from that of the activated nanoparticles; forming a complex and providing therapy to the patient by a method that administers the complex to the patient under 50 conditions to result in improved therapy to the patient by a method that administers the complex comprising a plurality of nanoparticles and/or quantum dots containing an agent targeting the nanoparticles and/or quantum dots to a site under conditions sufficient to permit accumulation of the 55 complex at the target site, provides an energy source at the target site to penetrate the tissue and controllably heat the nanoparticles and/or quantum dots and generate thermal energy to induce a photoacoustic signal or sound wave from the nanoparticles and/or quantum dots, uses a processor to 60 control the amount of thermal energy delivered at the desired temperature to the target site, records the temperature and photoacoustic signal or sound wave from the target site or from one or more multiple locations, and amplifies and processes the recorded photoacoustic signal or sound waves to 65 generate a computational tomographic image of the nanoparticles and/or quantum dots at the target site. The patient ben-

efit may be, e.g., personalized therapy, enhanced therapy such as enhanced cellular uptake, enhanced cellular delivery, enhanced therapy duration, enhanced therapy outcomes, combined or synergistic therapy effects, etc., enhanced theranostics capability of an agent, an improvement in a therapy modality, etc.

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For example, enhanced theranostics capability may be achieved by

administering targeted nanoparticles that carry a therapeutic agent, e.g., a medicament and/or a biologic, and are ligated with polymers, administering thermotherapy, assessing the patient's response to the thermotherapy to determine a qualitative and/or quantitative therapy change based at least in part on the patient's response, and changing patient therapy based on this assessment.

In one example of the general embodiment, a complex of a piezoelectric nanoparticle and a gene is administered to a patient, and an externally-positioned ultrasound source activates the complex to control cell polarization, capable of complex activation in the absence of light penetration. Activation may occur through, e.g., skin, soft tissue, or skull. The piezoelectric nanoparticles may be administered proximate a peripheral nerve and may contain nerve growth factor. When activated by an external ultrasound source, the result is localized electrical stimulation to at least one of a muscle, tendon, joint, or ligament.

In another example of the general embodiment, nanoparticles containing medications and/or biologics such as genes are administered for use in the method in combination with methods for weakening the cell membrane. This facilitates influx or delivery of the nanoparticle-contained medications and/or genes into the cell, enhancing delivery and thus enhancing patient therapy.

In another example of the general embodiment, any of microwaves, alternating magnets, radiofrequency waves, or focused ultrasound is applied in conjunction with low dose X-ray radiation therapy, resulting in synergistic thermal and radiation therapy.

In another example of the general embodiment, the nanoparticles administered are a combination of gold nanoparticles and magnetic nanoparticles. For example, the nanoparticles may contain a magnetic core and a gold shell, and the gold shell may be radioactive.

In another example of the general embodiment, a nanoparticle/gene complex is administered proximate to an olfactory nerve of a patient, and energy is applied to the complex under conditions sufficient to activate the nanoparticles of the complex to result in brain cell therapy. Neuronal stem cells may be administered with the nanoparticles to enhance or repair neural brain cell function or deficiency.

In another example of the general embodiment, the complex is controllably heated at the target site using photoacoustic energy to a temperature of about 40° C.-42° C. to perturb lipid cellular membranes resulting in enhanced penetration of a medicament carried by the complex.

In another example of the general embodiment, the targeted nanoparticles carrying pepsin, chymotrypsin, etc. are administered. This results in localized cell membrane perturbation due to enzymatic action, and/or an enhanced patient immune response due to chemotactic action. For example, a tumor cell having a perturbed cellular membrane initiates an immune response that is enhanced relative to an antibodygenerated immune response, resulting in enhanced tumor cell damage.

These and other aspects of the invention will become apparent to one skilled in the art in view of the following detailed description of the invention.

## BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a schematic diagram of one embodiment of the invention showing a probe for hyperthermally treating tissue and visualizing a dye in the target site.

FIG. 2 illustrates therapy of a patient with a choroidal fumor.

FIG. 3 illustrates electromagnet administration non-invasively, through the conjunctiva to the tumor.

FIGS. 4A and 4B illustrate one embodiment after place- 10 ment of the magnet for a tumor affecting the lower leg and with a horseshoe-shaped electromagnet configuration.

FIG. 5 illustrates one embodiment after placement of the magnet for a facial skin tumor.

FIG. 6 shows the relationship between the photoacoustic 15 system, the processor, and the energy delivery system.

## DETAILED DESCRIPTION OF THE INVENTION

The present invention is directed to a method and compo- 20 sition for hyperthermally treating tissue. In particular, the invention is directed to as method for heating tissue above a temperature effective to kill tissue cells or inhibit multiplication of cells below the protein denaturization temperature of the tissue.

The method of the invention introduces a composition into the bloodstream or other system of the body in a location to flow into or through a target site to be treated. In one embodiment, the composition is introduced into the lymphatic system. A heat source such as generated by radiation energy is 30 applied to the target site to heat the tissue in the target site for a time sufficient to hyperthermally treat the tissue and activate the composition. As used herein, the term "hyperthermal" refers to a temperature of the cell or tissue that kills or damages the cells without protein denaturization.

The composition may contain a temperature indicator that is able to provide a visual indication when a minimum or threshold temperature is attained that is sufficient to hyperthermally treat the tissue. It is a feature of the invention to thermally effective temperature and to provide a visual indication that a temperature of at least 41° C., and preferably at least 42° C. is attained. In one embodiment, the composition includes a second temperature indicator to provide a visual indication when a protein denaturization temperature is 45 attained thereby providing an indication that a maximum desired temperature is exceeded. The heat source can be applied to the tissue so that the composition provides an indication that a thermally effective temperature is attained that is below the protein denaturization temperature of the 50 tissue.

The methods and compositions have been described in detail in various permutations and combinations. For example, various nanoparticles have been described by each of types, components, complexes, and compositions. As 55 another example, various "cargo" that the nanoparticles can deliver to a target site have been described, including therapeutic agents, i.e., medications, and biological agents, e.g., genes, etc. As another example, various activation methods have been described, e.g., ultrasound frequency, radiofre- 60 quency, etc., have been described. As another example, various additional agents that perform one or more additional functions have been described, e.g., agents that enhance cellular uptake or delivery. As another example, the disclosed methods may be used with other methods to achieve more 65 lasting effects, enhanced effects, a broader spectrum of effects, etc. As another example, the disclosed methods may

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be customized to provide more efficient or otherwise improved modalities. All of these selections result in an enhanced hyperthermal therapy benefit to a patient in need thereof. In some embodiments, the patient receives a direct benefit by a personalized therapy result and/or a theranostics approach. In some embodiments, the patient receives an indirect benefits by an improved or enhanced modalities in application of the method.

By targeted and methodical selection among the combinations of each of the above selections, features, and embodiments, enhanced benefits may be achieved. These benefits may be directly realized by the patient in patient-specific personalized therapy that may be achieved in all patients, or in desired patients, or under desired conditions. These benefits may be indirectly realized by the patient in enhanced methodologies that permit improved modalities. These benefits may be both in patient-therapy outcomes and in modalityfacilitation outcomes.

As one example, the use of nanoparticles for delivering medications and/or genes, in combination with methods for weakening the cell membrane in order to facilitate influx or delivery of the nanoparticle-contained medications and/or genes into the cell, enhances delivery and thus enhances therapy. As another example, the combination of thermotherapy achieved by the application of any of microwaves, alternating magnets, radiofrequency waves, or focused ultrasound, in conjunction with low dose X-ray radiation therapy, results in synergistic therapy.

In one embodiment, the nanoparticles provide a process of diagnostic therapy for individual patients to assess reaction to therapy and adjust or tailor a therapeutic regimen based on the initial results. For example, administration of targeted nanoparticles carrying any medication (biological or non-biological agent), ligated with polymers, in combination with ther-35 motherapy, constitutes the basis of targeted drug delivery used for managing any disease processes, as the nanoparticles serve a theranostic capability, i.e., providing both diagnostic and therapeutic functions.

More specifically, and as only one non-limiting example, a provide a method of heating tissue in a target site to a hyper- 40 method of therapy by administering nanoparticles that are a combination of gold nanoparticles and magnetic nanoparticles, e.g., nanoparticles containing a magnetic core and a gold or a radioactive gold shell, targeted to a specific site, provides maximal therapeutic effect at low doses. Nanoparticles containing a gold shell or gold nanoparticles themselves may be radioactive or may enhance the effect of X-ray radiation.

One non-limiting example of a benefit achieved or realized by a modality improvement outcome is administering a piezoelectric nanoparticle/gene conjugate. Use of a piezoelectric nanoparticle/gene conjugate permits use of an externally positioned ultrasound to control cellular polarization where the surrounding medium is not sufficient clear to permit light penetration to achieve complex activation. This embodiment may be used through tissues such as skin, soft tissue such as muscle), hard tissue such as skull, etc. As one example, injecting piezoelectric nanoparticles proximate a peripheral nerve, with the nanoparticles additionally containing nerve growth factor, then activating these peripherallylocated nanoparticles permits a localized electrical stimulation using an external ultrasound source, providing therapy to muscles, tendons, joints, ligaments, etc. to improve skeletomuscular function. As another example, a plurality of nanoparticles carrying excitatory genes, e.g., opsin genes, may be delivered with piezoelectric nanoparticles such as zirconate titanate, perovskite-based nanoparticles, oxides, barium titanate, (poly)vinylidene fluoride (PVDF), and quantum

dots. Delivery may be to the brain, spinal cord, peripheral nerves, retina, heart, etc. Stimulation of the piezoelectric nanoparticles with extra-corporal ultrasound initiates an electrical response. If nanoparticles are present in, e.g., an implantable small diode, the system can initiate an on/off response as desired for diode laser light production, thereby controlling the pulsatile action potential response of the excitable cells carrying an opsin gene and rhodopsin membrane channels. This in turn initiates an on/off action potential in the membrane of the excitable or non-excitable cells, depending on the frequency of the applied light pulse. As another example, piezoelectric nanoparticles coated with a biocompatible molecule may be injected in the body close to peripheral nerves, resulting in nerves that can be directly stimulated by an externally located focused ultrasound probe. The piezoelectric nanoparticles convert the sound wave into an electric signal that propagates to adjacent tissues and nerves. The result is controlled stimulation of the peripheral nerves, e.g.,

Another non-limiting example of a benefit achieved or realized by a modality improvement outcome is the ability to activate brain cells via a nasal route. In this embodiment, after administration of a nanoparticle/gene complex close to the olfactory nerve, energy is applied to the complex under conditions sufficient to activate the nanoparticles of the complex. As an additional benefit, depending upon patient needs and a desired outcome, neuronal stem cells may be co-administered with the nanoparticles either as part of the complex or separate from the complex, and the method performed as 30 described. This is a unique way of controlling the brain action potential, and other brain cell functions.

to repair paresis, paralysis, spastic muscles, numbness, etc. 20

Another non-limiting example of a benefit achieved or realized by a modality improvement outcome is targeted hyperthermal therapy using targeted cellular penetration of 35 the nanoparticle complex. In this example, the use of photoacoustic technology enhances cellular penetration of the nanoparticle. As the nanoparticles are heated by an external energy source, they perturb or "melt" the cellular membrane phospholipid bilayer components at a temperature of about 40° 40 C.-42° C. The perturbed cellular membrane has increased permeability to the medicament carried by the nanoparticle complex. At temperatures exceeding about 42° C. to about 45° C., the cellular membrane is further damaged, creating an open pathway to the cell cytoplasm. Similarly, a non-thermal, 45 localized, cell membrane perturbation method involves the combination of targeted nanoparticles carrying enzymes such as pepsin, matrix metalloproteinase (MMP), chymotrypsin, etc. Such enzymes, conjugated to nanoparticles, are released at a tumor site at a desired temperature under control of a 50 photoacoustic technology unit. The nanoparticles deliver these enzymes that hydrolyze or otherwise degrade or dissolve membranes of cancer or other cells locally after the enzymes are released, thus enhancing delivery of drugs and/ or genes to these cells. Besides creating localized damage to 55 the cell membrane, these enzymes also are chemotactic, thus beneficially enhancing an immune response in the patient. In one embodiment, enzymes such as pepsin, alpha chymotrypsin, trypsin, MMPs, etc. effect lysis that can be therapeutic in patients with Alzheimer's disease by locally dissolving the 60 associated plaques and/or scar tissue. After application, other nanoparticles containing nanoparticles and medicaments are administered to block the enzymes and prevent excessive inflammation at the tumor site.

When the cellular membrane perturbed by the method is a 65 tumor cell, these damaged tumor cells initiate an immune response that is enhanced, compared to other antibodies,

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attracting monocytes as well as other leukocytes and other killer cells to eliminate the tumor.

In one example, the targeted nanoparticles are heated with a source of energy such as radiofrequency waves, microwaves, focused ultrasound, electromagnetic radiation, alternating magnetic field, etc. The nanoparticles expand and produce a photoacoustic signal that can be acquired by one or multiple receivers attached to the patient's body. The sound waves are recorded and imaged in two or three dimensions with software as a photoacoustic image, indicating the exact location and temperature of the nanoparticles at that location. These images are overlapped with a processor and software on simultaneously obtained images using X-ray, MRI, fMRI, ultrasound, PET-scan, CT-scan etc. for topographical localization and follow up. The use of nanomaterial such as gold nanoparticles, carbon nanotubes, magnetic particles, functionalized quantum dots, etc. offer multiplexing capability for simultaneous measurement of multiple cancer biomarkers and enhanced molecular imaging of the cancer while simultaneously providing therapy to the patient.

In one example, smaller sized ferric oxide nanoparticles of 1 nm-10 nm are selected to heat the cell membrane more rapidly than larger sized dendrimer nanoparticles or liposomes of size 2 nm-200 nm. The smaller nanoparticles absorb more thermal energy than larger nanoparticles due their larger collective surface area. It is known that the thermal energy absorption is related to the size of the nanoparticles. Thus, for example, ferric oxide nanoparticles, in comparison to, e.g., a dendrimer, heat more rapidly because of their molecular composition. Heating ferric oxide nanoparticles to a temperature of, e.g. at least 45° C., damages the cell membrane, rendering it more deformable and "leaky", while the larger dendrimer and liposome have released their contained medicaments and/or biologics, from their temperature-sensitive polymeric coating. At the same time, the damaged cell membranes of the targeted cells permit ease of influx of other nanoparticles containing medicaments and/or biologics.

The use of nanoparticles for enhanced delivery of a medicament and/or a biologic, at a defined temperature and location in the body, under the control of a photoacoustic unit, has never been reported.

The biological agent may be an inhibitory gene, e.g., sRNA, microRNA, etc. The nanoparticle complex containing an inhibitory gene is administered to reduce the tumor-promoting potential of cancer cells, or to reprogram the cancer cells toward their original cell type, thus differentiating the cancer cells back to a non-pathologic phenotype. It will be appreciated that use of the method to deliver inhibitory genes may be combined with delivery of other biologic and/or non-biologic medicaments. Such medicaments include, but are not limited to, antineoplastic agents, antibodies, therapeutic drugs, immunotherapy agent, vaccines, etc. These medicaments are also conjugated or otherwise associated with the nanoparticles as part of the nanoparticle complex administered to the patient.

It is known that transcription factors initiate production of a new protein in cells. Transcription factors bind to a specific DNA region to prompt gene transcription; creating from the DNA template an RNA template for a new protein. Changing the activity of these factors could influence creation of proteins: e.g., increase production of proteins that suppress tumors or reduce inflammation, and even reprogram adult cells into immature cells or new cell types. Such activity changes are not long lasting, unlike gene therapy, thus making it desirable for treatment of tumors, inflammatory disease, atherosclerotic disease, liver disease, etc. However it is difficult to deliver transcription factors because they are large

proteins that, unprotected by DNA, are subject to lysosomal degradation after entering the cytoplasm. Such large proteins need to be protected by "wrapping" them with DNA that cannot be destroyed in the acidic milieu of cell lysosomes. In one embodiment, the desired transcription factor is protected by DNA or aptamer conjugated-targeted nanoparticles for thermal assisted gene and/or medicament delivery inside the cell after their systemic or local injection.

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In one embodiment of the inventive method, not only is the DNA of a cell nucleus modified, but abnormal DNA that can 10 cause mitochondrial DNA disease is damaged. In this embodiment, the targeted nanoparticles are delivered to release a DNA-endonuclease or a CRISPR/Cas9 complex in the cells. The endonuclease and/or the CRISPER/Cas9 complex then seek DNA containing a specific mutation in the 15 mitochondria or the nucleus. The cleavage functions of the endonuclease and/or CRISRP/Cas9 complex destroy the disease-producing DNA by excising it or replacing it with a normal DNA segment, as known in the art. In another embodiment, one delivers stimulating genes, such as an opsin 20 family gene, to repair or replace a defective gene in the nucleus or mitochondria of retinal cells, CNS cells, neuronal cells, glial cells, spinal cord cells, peripheral nerve cells, etc., in a nanoparticle conjugated CRISPR/Cas9 complex. In another embodiment, a plurality of nanoparticles is adminis- 25 tered to an excitable cell in, e.g., retina, CNS, spinal cord, peripheral nerves, heart, etc., and uses photoacoustic technology and thermal energy to control delivery of multiple genes and at the same time effect repair of a genetic disease along with enolase or CRISPR DNA while delivering an additional 30 stimulating gene such as an opsin gene, etc. The cells are stimulated with light pulses of wave lengths of, e.g., 400 nm-infra red, and mid-infrared. This embodiment treats or ameliorates diseases such as epilepsy, depression, pain, Alzheimer's disease, Parkinson's disease, spinal cord inju- 35 ries, heart failure, arrhythmia, etc. In one embodiment, CRISPR DNA conjugated nanoparticles are administered locally (e.g., eye) or systemically. Once CRISPR DNA enters the cell, it is transcribed into a nuclease that scans the host DNA for a target sequence. The nuclease cuts the host DNA 40 at the target. With one cut, CRISPR can insert a new custom gene sequence; with two cuts, two CRISPRs can excise the DNA, providing the host with a new DNA.

It will be appreciated that use of the method to deliver inhibitory genes may be combined with delivery of other 45 biologic and/or non-biologic medicaments. Such medicaments include, but are not limited to, antineoplastic agents, antibodies, therapeutic drugs, immunotherapy agent, vaccines, etc. These medicaments are also conjugated or otherwise associated with the nanoparticles as part of the nanoparticle complex administered to the patient.

Vaccines can be used simultaneously with nanoparticles for immunotherapy or other therapeutic indications. For example, in one embodiment, nanoparticles deliver inhibitors to the target cells. The inhibitors include, but are not limited 55 to, CTLA-4 (Ipilumab), PD-1, B7-H1 producing INFgamma, and other checkpoint inhibitors or virocytes for the cancer therapy. By simultaneously damaging the tumor cell membrane using thermotherapy, more tumor antigens are released, which attract monocytes, dendritic cells, killer 60 T-cells, macrophages, etc., contributing to elimination of the cancer cells. In each case, the vaccine can trigger an increase in T-cells that could recognize the patient's cancer cell. As non-limiting examples, the method can use polio vaccine for therapy of glioblastoma, AIDS vaccine, Ebola vaccine, and 65 herpes or papilloma vaccine for different cancers. Customized vaccines containing synthetic genetic sequences can be

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used. In one embodiment, the combination of vaccine therapy and thermotherapy is applied to treat or prevent an infectious disease while simultaneously delivering multiple antibiotics, antivirals, antifungals, vaccines, etc. in treatment of therapy resistant organisms.

In one embodiment, cerium oxide nanoparticles are used. Molecular substrates are conjugated with the cerium oxide nanoparticles, and the nanoparticles are administered locally, systemically, or in vitro in a stem cell culture to support stem cell survival and function. This embodiment controls production of intra- and extracellular oxygen species, and encourages stem cell proliferation and differentiation. Neuronal, muscular, mesenchymal, epidermal, and other stem cells can be used. Other nanoparticles such as dendrimers or liposomes can be used, when administration simultaneously provides nerve growth factor or other medicaments to facilitate neuronal growth.

In one embodiment, nanoparticles are conjugated with a virus such as adeno-associated virus (AAV) for gene delivery. This embodiment enhances success of viral gene delivery by creating, initially, a weakened area in in the thermally-damaged targeted cell membrane. The gene can be delivered in a localized organ, e.g., eye, CNS, heart, spinal cord, peripheral nerves; the gene can be inhaled; the gene can be locally injected or systemically administered for treatment of tumors, infection diseases, degenerative diseases (e.g., Alzheimer's disease, Parkinson's disease), spinal cord injuries, and peripheral nerve injuries; the gene can be introduced in vitro to cells in culture to manipulate DNA of stem cells using a CRISPR/cas9 complex or to enhance function prior to patient administration.

In one embodiment, the nanoparticles are used in conjunction with an electrical current to affect cell membrane polarization. Constant hyperpolarization and depolarization affect membrane sodium, potassium, and other membrane channels, rendering the membrane more permeable to agents, e.g., chemotherapeutic agents. Alternatively, constant hyperpolarization and depolarization may have an additive damaging effect with thermal radiation, X-ray radiation, or other types of radiation on cancer cells. This concept may be used in cancer therapy for local damage of cancer cells. Targeted coated piezoelectric nanoparticles and/or quantum dots may be injected locally or systemically to seek cancer cells, to which they attach or are internalized. They may then be subjected to ultrasound stimulation applied from a remote site to induce an electrical current in the nanoparticles and/or quantum dots, which is transmitted to tumor cells, render the tumor cells more vulnerable to other forms of therapy.

In one embodiment of the invention, the method introduces a composition to a target site, where the composition includes a fluorescent dye that is encapsulated in a heat sensitive particle, such as a liposome. The dye is a fluorescent dye that can be excited to fluoresce and be observed or visualized by the operator. Preferably, the heat sensitive liposomes are formed to rupture or release the fluorescent dye at a temperature at least equal to the temperature necessary to kill cells in the tissue and at a temperature below the protein denaturization temperature. The composition containing the heat sensitive liposomes encapsulating the fluorescent dye is introduced into the bloodstream to flow to or through the target site. The amount of the liposome composition is introduced in an amount effective to be released in or near the target site and to excited and visualized by the exciting light source and the visualizing device. The composition containing the dye can be injected in a single dose into the bloodstream or injected continuously to supply a continuous flow of the composition through the target site. The amount of the composition intro-

duced can vary depending on the target site and the length of time that the dye is to be excited. A light or energy source is continuously applied to the target site to excite the dye and to cause the dye to fluoresce when released from the liposomes. An imaging device is used to capture the fluorescing light 5 from the dye to provide a visual indication that the dye is released. The release temperature of the liposomes are selected to release the dye at a predetermined temperature so that when the dye is fluoresced and visualized, the visualization provides the operator with an indication that the release temperature in the target site has been attained. In one embodiment, the liposome composition is injected into the blood stream so that the composition is able to provide a continuous supply of the dye for fluorescing during the hyperthermal treatment. In this manner the operator is provided 15 with a continuous indication that a sufficient temperature is being maintained.

The method of the invention is primarily directed to a method of heating tissue and cells in the tissue of an animal, particularly a human patient, at least to the temperature suf- 20 ficient to kill or damage the cells. Cell death or cell damage is known to occur when the tissue cells are heated to a temperature of about 5° C. above the normal body temperature of 37° C. Therefore, the method of the invention heats the cells in the tissue to a temperature of about 41° C., and preferably at least 25 42° C. for a time sufficient to kill or damage the cells. Preferably, the heat source is applied to minimize unnecessary damage to the surrounding cells and tissue.

In one embodiment of the invention, the tissue is heated to a temperature of at least 41° C. and preferably in the range of 30 at least about 42° C. to about 50° C. Heating the tissue to at least 42° C. ensures that a sufficient temperature is obtained to thermally treat the tissue and the cells effectively. Preferably, the tissue is heated to a temperature below the protein denaturization temperature of the tissue. Protein denaturization 35 begins to occur at about 50° C. to 51° C. and occurs rapidly at temperatures of about 60° C. Preferably, the tissue is heated to a temperature of less than 60° C. and more preferably to a temperature of about 50° C. or less.

In one preferred embodiment, the tissue and the cells are 40 heated to a temperature of about 47° C. to about 49° C. for a time sufficient to kill or damage the cells without protein denaturization. The length of time that the tissue is heated will depend on the location of the target site, the size and dimensions of the target site, the desired depth of penetration of the 45 heat and the desired extent of thermal treatment- or damage of the tissue and cells in the target site. Typically, the heat source is applied for several minutes. In one embodiment, the heat source is applied for about 1 to 15 minutes, and typically about 5 to 10 minutes.

The heat source can be applied to a variety of the areas in the body where the hyperthermal treatment is desired. The target site can be tumors, organs, muscles and soft tissue. Examples of a target site include blood vessels and arteries, able for hyperthermally treating the epithelial cells on the lens of the eye after cataract surgery. Other target sites include the retina and the choroid.

In other embodiments the target site may be cell components of various organs. The organs may be healthy or may 60 contain tumors, either malignant or benign. The following are representative, not limiting, examples of cell components on which the inventive method may be applied: tumors of the central nervous system (CNS), various layers of skin and its underlying support structures, intestinal tract, kidney, urinary tract, female and male reproductive system organs, bone including bone marrow, circulatory system components

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including the heart, blood vessels, and circulating malignant cells, the lymphatic system including lymph nodes and vessels, and the respiratory system.

In one embodiment, the compositions including, e.g., gold or ferromagnetic nanoparticles, as described below, are injected through the nipple. The injected composition travels through the duct leading down from the nipple ending in glands, i.e., acini aggregated into lobules. This mode of injection may also introduce the composition into the lymphatic system, particularly if injected into the breast stroma. This embodiment may be used to treat breast cancer, as well as cancers in the associated lymphatic tissue. In addition, this embodiment may be used as a prophylactic treatment to obliterate or substantially reduce the breast glands' secretary epithelium in patients who exhibit genetic predisposition to breast cancer.

In various embodiments, the described compositions are introduced through any accessible cavity, such as oral, respiratory, or genitourinary cavities. The compositions may be introduced by needle injection or via a catheter. In one embodiment, the described methods may be used to treat cervical or bladder cancer, including early stages of the dis-

In one embodiment, cell proliferation in tumors that would otherwise be excised from a body surface or removed from an internal body site are treated by the inventive method. Treatments involves both amelioration and/or tumor debulking, i.e., reducing the tumor mass. In one embodiment, the inventive method is used to augment surgical removal of a tumor. Tumors that are relatively large are often surgically removed, even if the tumor is determined to be non-malignant, because of the space-occupying nature of the tumor and/or the stress on other organs. For examples, stress on adjacent organs, such as liver and kidneys, can result in the potential for hepatic and nephrotic complications.

During surgical tumor excision or removal, it is important to ensure the tumor is completely removed, yet the tumor margins are difficult to recognize and determine. Typically, complete tumor removal requires multiple biopsies, during the surgical procedure itself and while the patient is under anesthesia, from the edges of the tumors. The biopsies are histologically or otherwise evaluated in real time by a pathologist. This very time consuming task prolongs surgery, may put the patient at increased risk, and definitive pathology results may take hours to obtain. Often, even after careful excision of the tumor, subsequent histologic studies of the entire lesion show that not all margins or edges are free of tumor tissues, resulting in patient anxiety, increased cost for subsequent surgery, and heightened risk of post-surgical complications. Such uncertainty of complete tumor and/or cancer cell removal makes subsequent medical treatment difficult to determine, e.g., whether to remove lymph nodes located in the area of the tumor, etc.

Malignant tumors have a preferred location in the body esophagus and eyes. In one embodiment, the method is suit- 55 from which they grow from the initial mother cell. They then metastasize in other organs. The initial tumors are seen in any organ of the body (e.g., eye, brain, skin from which retinoblastomas, gliomas, melanomas occur; also prostate, colon and other intestinal tumors, pancreas, thyroid, etc. Tumors can grow from any type of tissue (e.g., ectoderm, mesoderm, endoderm, neuro-ectoderm). Using the eye as one example, melanomas can originate from the uveal tract, conjunctiva, orbit, etc. Melanomas of the choroid, ciliary body, and iris grow very slowly, in contrast to cutaneous melanomas, and remain in place before metastasizing. More rapidly growing melanomas preferentially metastasize in the liver, but are not limited and may metastasize to any organ. Prostate cancer

preferentially metastasizes to bone. Colon cancer preferentially metastasizes to liver. Some tumors, such as basal cell carcinoma, gliomas, meningioma, etc. grow locally but damage vital organs.

It is thus desirable to treat the early diagnosed tumors of locally and with a more intense therapy such as using ferromagnetic nanoparticles, with or without a single or combination of drugs, that can be attracted to a locally positioned magnet, either external or internal. Placement of a naturally occurring magnet or electromagnet, with a stable magnetic field, adjacent to or over the tumor area attracts and concentrates the circulating ferromagnetic particles to that specific location during their circulation in the blood after administration (e.g., injection) into the circulatory system, body cavity, etc., or at the site of administration, e.g., when topically applied to the skin, mucosa, etc.

In this embodiment, using a magnet to localize ferromagnetic particles at a tumor or lesion site, or even using whole body therapy, thermotherapy is controllably applied to a patient to treat a localized tumor and metastasized lesions.

Activatable cell-penetrating peptides (ACPPs) that are labeled with fluorescent polycationic cell-penetrating peptide (CPP) coupled by a cleavable linker to a neutralizing peptide have been developed and utilized to visualize tumors during surgery. Similarly there are other tumor specific biomarkers 25 that are known to one skilled in the art, used for breast tumor, genitourinary tumors, lung tumors, head and neck tumors, gastrointestinal tumors, brain tumors, etc., as only one nonlimiting example, disclosed at http://www.medscape.org/ viewarticle/725989. ACPP conjugated to dendrimers (ACP- 30 PDs) and gadolinium chelates can allow MRI visualization of whole body tumors, permitting thermotherapy if the magnetic or gadolinium nanoparticles are labeled with ACPPD. However, gross observation of a labeled tumor does not guarantee that all the tumor margins will be visible or will 35 removed. One reason is that tumor margins may contain sparse numbers of dispersed cancerous cells. Another reason is that cancerous cells may be embedded more deeply in the tissue and thus not amenable to labeling (e.g., staining) and/or visualization. Thus, while such a method helps the surgeon 40 during a surgical procedure, it does not eliminate the need for a biopsy from the tumor margin or edge.

The inventive method provides safe surgical debulking of a large tumor, while concomitantly providing that the tumor margins or edges are free of viable cancerous cells, or that 45 such cancerous cells are non-viable. The method removes and treats non-malignant tumors, damaged cells such as nerve cells, or normal cells involved in or affected by adverse immune reactions in organ transplant procedures.

One embodiment is a composition of a nanoparticle that is 50 conjugated to an ACPP. The nanoparticle may be cleavably conjugated to the ACPP by, e.g., a linker (e.g., at least ethylene glycol moiety or a (poly)ethylene glycol moiety). In one embodiment, the nanoparticle is labeled with a label such as a fluorescent moiety, chemiluminescent moiety, etc. In one 55 embodiment, the ACPP is labeled with a polycationic cellpenetrating peptide (CPP).

In one embodiment of the method, the above-described composition is prepared with any biocompatible excipient and is administered to a patient. In one embodiment, a regimen of low dose medication is prescribed to enhance the therapeutic effect of the inventive method including but not limited to at least one antiproliferative agent, e.g., cisplatin, carboplatin, tetraplatin, iproplatin, adriamycin, mitomycin C, actinomycin, ansamitocin and its derivatives, bleomycin, 65 Ara-C, daunomycin, metabolic antagonists such as 5-FU, methotrexate, isobutyl 5-fluoro-6-E-furfurylideneamino-xy-

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1,2,3,4,5,6 hexahydro-2,4-dioxopyrimidine-5-carboxylate, melpharan, mitoxantrone, lymphokines, etc.

One embodiment provides a method of drug delivery and thermal damage to a target in a patient in need thereof. This embodiment provides four ranges of thermotherapy at the target site. The target may be a particular site in the body, e.g., a tumor site. The target may be an organism, e.g., a bacterial target. Each temperature range results in a specific effect, as subsequently described. The overall result of this embodiment is specific drug delivery and specific thermal treatment of the target.

This embodiment is based on the knowledge that pores in bacterial membrane increase specifically during bacterial cell growth. Pores can serve as a conduit for transfer of genes and plasmid DNA from other bacteria that can effect cell survival, e.g., render the bacteria drug-resistant. Bacteria can be engineered to produce proteins, e.g., insulin, by transmitting the genes encoding such proteins to the bacteria interior. One skilled in the art appreciates that the pore size and number of pores increase with a cell temperature increase in the range of about 2° C. to about 28° C. At a temperature of greater than about 40° C., bacteria become very vulnerable, e.g., they become particularly sensitive to drugs and frequently die. While the same mechanisms take place in mammalian cells, mammalian uptake occurs more by pinocytosis.

One embodiment is a method to target therapy in a patient in need thereof. An antibody-coated and/or drug-containing nanoparticle composition is administered to the patient, then stages of increasing temperature thermotherapy at the target containing the antibody specific and/or drug-coated nanoparticles are performed. In one embodiment, the target is imaged during the temperature increases. The thermotherapy is administered under conditions (e.g., temperatures, durations, etc.) sufficient to result in targeted therapy in the patient. The inventive method limits thermotherapy to a specific target, e.g., a tumor, bacteria, etc. and does not substantially affect normal cells proximate this target. In one embodiment, the method is performed on a patient receiving therapy, e.g., chemotherapy, radiation therapy, anti-vascular endothelial growth-factor therapy, and/or steroid therapy. In one embodiment, the anti-tumor antibody-coated or -containing nanoparticles further comprise a thermosensitive polymer which releases the drug contained in the nanoparticle by dissociation of the polymer when the thermosensitive temperature is attained. Thermosensitive polymers include chitosan-poly (N-isopropylacrylamide), smart polymers, poly(N-isopropylacrylamide (PNIPAM), poloxamers, poloxamines, and/or acid (PMA) polymers modified with thiol groups (PMA<sub>SH</sub>), polymers containing polyorthoesters, polyglycolic acid, polyether esters, liposomes, polyalkylcyanoacrylates, polycarbonates, polyhydroxyalkanoates, polyhydroxy-acids, poly- $(\epsilon$ -caprolactone), polyamino-acids, polysaccharides, polysaccharide-(meth)acrylate derivatives, isopropyl cellulose, hydroxyalkyl celluloses, cellulose acetate butyrate, starch, starch alkyl ethers, collagen, alginate, chitosan, polyvinyl alcohol, gelatin, ionically crosslinked multi-functional amines or linear copolymers, graft copolymers or block copolymers of these polymers, or dendrimers formed from said these polymers.

In use, in one embodiment, the composition is administered to the patient with four subsequent staged increases in temperature. In one embodiment, the temperature at the target is first increased to result in drug release from the nanoparticles. These conditions typically occur with a temperature at the target of at least 35° C. to 43° C., which is provided for about 0.5 minutes to about 20 minutes. The temperature at the target is then increased to result in thermotherapy. These

conditions typically occur with a temperature at the target of 43° C. to 50° C., which is provided for about 1 min to about 15 min. The temperature at the target is then increased to result in expansion of the gases that are dissolved in cellular fluids and concomitant expansion of the membrane, cytoplasm, and/or nuclear pores. Such expansion results in mechanical distension of the cell membrane and/or nuclear membrane with subsequent expansion of the membrane pores and/or channels allowing essentially unrestricted drug into the cell. These conditions typically occur with a temperature from 50° C. to 60° C., which is provided for about 1 min to about 10 min. The temperature at the target is then increased to result in water evaporation at the target, causing cumulative damage, i.e., essentially unrestricted drug flow combined with protein denaturation and water evaporation. These con- 15 ditions typically occur with a temperature greater than 60° C. (i.e., a temperature that kills normal cells) and up to 100° C., which is provided for a duration less than one second to a few

As one example, in one embodiment, the composition is 20 administered to the patient, then the temperature increase at the target is limited to less than 60° C. The tumor cells and/or bacterial cells, which are rapidly dividing and hence have increased susceptibility to increased temperature, are killed.

Thermotherapy may be provided by, e.g., electromagnetic 25 radiation, ultrasound energy, or an alternating magnetic field. Imaging of the target during the temperature increases can control thermotherapy by connecting the photoacoustic system, which controls the heat generating systems, to the electromagnetic radiation-creating instrument, or to a reversible 30 magnet, or another ultrasound system generating a focused ultrasound beam. This connection may be, e.g., by a processor. The operator instructs the system to achieve the desired temperature, or provides an algorithm for the system to follow to create the desired result, using specified temperatures and 35 exposure durations. Imaging may be thermal imaging, photoacoustic imaging, X-ray imaging, optical coherence tomography, ultrasound imaging, fluorescence imaging, chemiluminescent imaging, positron imaging, surface enhanced Raman spectroscopy, and/or magnetic resonance imaging 40 (MRI). In one embodiment, imaging is by MRI.

The composition administered may contain magnetic, diamagnetic, ferromagnetic, and/or paramagnetic nanoparticles. The composition administered may contain gold nanoparticles, diamond nanoparticles, platinum nanoparticles, and/or 45 carbon nanoparticle.

The targeted nanoparticles, that contain antibody as well as drug in or on the particles, are administered to a patient in need thereof. The staged and selective temperature increases administered at the target deliver drug from the nanoparticle 50 to the cell interior, with heat increasing metabolic need of the target (e.g., tumor cells or bacteria). Thermal therapy is staged as follows: a first temperature at the target site of at least 35° C. to 43° C., thereafter a second temperature at the target site of from 43° C. to 50° C., thereafter a third tempera- 55 ture at the target site of from 50° C. to 60° C., and thereafter a fourth temperature at the target site of greater than 60° C. The duration of the first stage is from 0.5 min to 20 min. The duration of the second stage is from 1 min to 15 min. The duration of the third stage is from 1 min to 10 min. The 60 duration of the fourth stage is from less than 1 second to a few minutes. In embodiments, a stage of thermotherapy can be removed. For example, the method may be used in a patient with a tumor that is drug resistant, but the tumor will not be able to develop resistance to the increased temperature.

The inventive thermotherapy method results in specific drug delivery and thermal damage to target cells. The method 36

delivers and releases drug from the nanoparticles to the cell interior. The increased temperature increases the tumor cell/bacteria metabolic needs. Without being bound by a specific theory, the result is cell damage and/or death by drug, by heat, or by the combination of drug and heat.

The staged, selective temperature increase in the nanoparticle can also expand the gas that are dissolved in the cell fluid, creating a mechanical stress/damage on the cells/bacteria membrane or internal metabolic machinery. The internal gas expansion distends the wall of the cell/bacteria and expands the pore size of the cell/bacteria membranes. This in turn causes almost free flow of drug to the cell interior, resulting in cell death. Increasing the temperature, while imaging, beyond 60° C. can also cause not only protein denaturation in the cell, but also cause the water molecules to evaporate. This further increases the cell damage leading to cell death. The different stages can be controlled by the imaging technology described controlling the thermal effect of the nanoparticles. The processes occur more rapidly and more readily (faster and sooner) around the nanoparticles than around the surrounding tissue because of the nanoparticles' small size.

Administration may be by any route and at body site. For example, the composition may be administered by an intrathecal, intravenous, intraocular, etc. route. The composition may be administered into a body cavity (e.g., eye, bladder). The composition may be administered into the cerebrospinal fluid. The composition may be applied topically, e.g., to an external or mucosal lesion. In one embodiment, the composition is intravenously administered. The composition may be administered locally in or near the tumor site, or in or at any body site.

The composition contains a nanoparticle conjugated to an activatable cell penetrating peptide (ACPP), which forms an antitumor-ACPP-nanoparticle-cell complex at the tumor tissue site, or target site (e.g., liver). The target site of the patient is exposed to an energy source under conditions sufficient to heat the nanoparticles in the antitumor-ACPP-nanoparticle-cell complex to a temperature that is sufficient to measure an acoustic response produced by the nanoparticle. Exemplary durations are from less than one second to 15 minutes. The treatment can be repeated as needed, e.g., within weeks, determined by MRI, CT, ultrasound, PET scans, etc.

In one embodiment, the temperature to measure an acoustic response produced by the nanoparticle, indicating damage to the target cells, ranges from about 40° C. to about 60° C. inclusive. A lower threshold temperature of 39° C. may be used with drug delivery to break up the medication-nanoparticle complex at the tumor site. In one embodiment, the temperature to measure an acoustic response produced by the nanoparticle, indicating damage to the target cells, ranges from about 42° C. to about 45° C. inclusive, or from about 45° C. to about 50° C. inclusive, or from about 50° C. to about 58° C. inclusive. In one embodiment, for superficially located or otherwise externally accessible tumors, or for treating tumor margins after surgical excision or debulking, a temperature >58° C. up to 60° C. is acceptable if the target is away, e.g. distal or remote, from a vital organ. Such hyperthermal therapy damages or kills the tumor cells, referred to as tumor de-bulking.

In one embodiment of the method, the tumor is surgically excised or removed prior to or concomitantly with administering the composition and performing the method. The surgery thus defines the lesion that is a target area for treatment by the method.

The method may be used on targets or lesions in and/or on any body site. Examples include, but are not limited to, skin, mucosa, organs and tissues of the digestive, genitourinary,

nervous (CNS and/or peripheral nerves), respiratory, circulatory, lymphatic, and other systems. As one example, a tumor may be located in and/or on a mucosal lining or skin. As one example, cancerous blood cells may be treated by the inventive method, e.g., leukemia or lymphoma. In one embodiment, intravenous administration of the composition permits therapy to circulating blood or other cells (e.g., immune cells). With surgical tumor excision, the composition can be intravenously injected prior to, after, or during surgery.

In one embodiment, use of the method to eliminate specific immune cells effectively reduces or eliminates the need for immunosuppressive drugs in cases of organ transplant. In such an embodiment, nanoparticles are labeled with specific binding molecules that bind to specific surface receptors of these immune activated cells; the method proceeds as described. This embodiment minimizes or eliminates the need to expose the entire body to potentially damaging radiation, chemotherapy, and/or immunosuppressant medications, thus minimizing or eliminating their damaging side effects.

In one embodiment, use of the method combines hyper-thermal treatment, as disclosed herein, with immunotherapy. The resultant immune-thermal therapy advantageously uses synergy from the combined approaches, specifically, (i) a patient's inherent response against a non-self substance, in 25 this case a tumor cell, to destroy the tumor cell, and (ii) localized thermal therapy possible by using the inventive method with targeted directed nanoparticles.

Thermal therapy, also termed hyperthermal therapy, is completely described, disclosed, and enabled herein. All theraple mal therapy embodiments may be used in the inventive method

In embodiments, thermal therapy may be performed simultaneously with, or substantially simultaneously with, immunotherapy. In embodiments, thermal therapy may be performed prior to immunotherapy. In embodiments, thermal therapy may be performed subsequent to immunotherapy. Because thermal therapy is targeted, involves thermal treatment without destroying proteins, and spares non-targeted tissue, it is generally relatively safer and less invasive than 40 chemotherapy and/or radiation therapy.

Targeted therapy includes, but is not limited to, anti-tumor antibodies, other specific binding compounds known in the art such as biotin with streptavidin, specific receptors and receptor agonists and/or antagonists, etc. Targeted therapy 45 may result when the nanoparticles contain and/or are coated with one part of the associated pair and the tumor itself and/or an area proximate to the tumor contains the complementary part of the associated pair. As another example of targeted therapy, the nanoparticles may be completely or partially on, 50 associated with, and/or contained in a virus to more specifically target therapy. As one example and without limitation, the nanoparticle may be targeted to a site of a tumor that has or may have a viral association. Examples include, but are not limited to, papillomaviruses, polyoma viruses, Epstein-Barr 55 virus (EBV), human herpes virus 8 (HHV-8), hepatitis B virus, and human T-cell lymphotropic virus-1 (HTLV-1). As another example, the nanoparticle may be targeted to, incorporated in, or otherwise located at a tumor cell, e.g., membrane viral capsids and/or envelopes, using a modified-virus 60 and/or adenoma-associated virus (AAV) either alone or in combination with another therapy, e.g., gene therapy. As another example, the nanoparticle may be completely or partially on, associated with, and/or contained in a tumoricidal virus or a virus rendered tumoricidal to provide additional 65 therapy, e.g., a modified herpes simplex virus selectively targeting malignant cells.

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Immunotherapy is broadly defined as therapy to a patient that involves modulating at least some part of the immune system, in whole or in part, and whether the immune system involvement stimulates, modulates, suppresses, ameliorates, enhances, treats, etc. the patient. The immunotherapy thus may encompass either activation of the patient's own immune system or suppression of the patient's own immune system, in whole or, more likely, in part. Immunotherapy is used to treat neoplastic disease, infectious diseases, inflammatory disease, degenerative disease, etc.

Immunotherapy may provide one or more exogenous compounds that may be natural in whole or in part, recombinant in whole or in part, and/or synthetic in whole or in part. The type(s) of compounds may be diverse, e.g., small molecules, biologics, enzymes, hormones, protein or peptide factors, gene modulating agents, etc. Examples include, but are not limited to, antibodies including polyclonal and monoclonal antibodies, growth factors, integrins, interleukins, cytokines, chemokines, interferons, membrane-coupled receptors, receptor agonists, receptor antagonists, oligonucleotides, mRNA silencers, siRNA, steroids, etc. Upon administration to a patient, an immunotherapy compound acts on the patient's own immune system in one or more of the ways previously described, e.g., activation, suppression, enhancement, etc. The exogenous compound(s) may act directly or indirectly to provide an outcome. Thus the compound(s) may have a substantially immediate effect, or may have an attenuated or non-immediate effect.

Exemplary immunotherapy compounds include, but are not limited to, the following: polypeptides comprising MHC I and/or MHC II; glycoproteins; APC protein markers; dendritic cell maturation markers; polymers either with or without an lipid group, acrylic polymers, copolymers of lactic acid and glycolic acid; protein, peptide, small molecule, and/ or carbohydrate B cell antigens including toxins; carbohydrate targeting moieties; lipid targeting moieties; vaccines including the breast cancer HER2/neu peptide, cervical cancer HPV vaccine, colorectal cancer vaccine including vaccines against CEA protein, kidney cancer vaccine, lymphoma vaccine, lung cancer vaccine include BLP25 (STIMU-VAX®), pancreatic cancer vaccine including GVAX and ipilimumab (YERVOY®), prostate cancer vaccine including prostate-specific antigen (PSA) and prostate-specific membrane antigen (PSMA) and PROSTVAC-VF); Toll-Like Receptor (TLR)-1 to 10, TLR agonist; toxins; biologics including but not limited to the following: the farnesyltransferase inhibitor tipifarnib; farnesyl and geranylgeranyl transferase inhibitors; the multikinase inhibitor sorafenib combined with tipifarnib; and bortezomib and tipifarnib.

Exemplary markers to evaluate therapy include, but are not limited to, the following: alpha-fetoprotein (AFP) for liver and germ cell tumors; beta-2-microglobulin (B2M); betahuman chorionic gonadotropin (beta-hCG); BCR-ABL fusion gene; BRAF mutation V600E; CA15-3/CA27.29 in blood for breast cancer; CA19-9 for pancreatic cancer, gallbladder cancer, bile duct cancer, and gastric cancer; BCA-125 in blood for ovarian cancer; calcitonin in blood for medullary thyroid cancer; carcinoembryonic antigen (CEA) in blood for colorectal and breast cancer; CD20 in blood for non-Hodgkin lymphoma; chromogranin A (CgA) in blood for neuroendocrine tumors; chromosomes 3, 7, 17, and 9p21 in urine for bladder cancer; cytokeratin fragments 21-1 in blood for lung cancer; EGFR mutation analysis in tumors (estrogen receptor (ER)/progesterone receptor (PR)) for non-small cell lung cancer and breast cancer; fibrin/fibrinogen in urine for bladder cancer; HE4 in blood for ovarian cancer; HER2/neu in tumor immunoglobulins for breast, gastric, and esophageal

cancer, and in blood and urine for multiple myeloma and Waldenstrom macroglobulinemia; KIT in tumors for gastrointestinal stromal tumor and mucosal melanoma; KRAS mutation analysis in tumor for colorectal and non-small cell lung cancer; lactate dehydrogenase in blood for germ cell tumors; nuclear matrix protein 22 in urine for bladder cancer; prostate-specific antigen (PSA) in blood for prostate cancer: thyroglobulin in tumor for thyroid cancer; urokinase plasminogen activator (uPA) and plasminogen activator inhibitor (PAI-1) in tumor for breast cancer; 5-protein signature (Ova1) in blood for ovarian cancer; 21-gene signature (Oncotype DX) in tumor to evaluate risk of recurrence in breast cancer; 70-gene signature (Mammaprint) in tumor for breast cancer. Generally, a decrease from a patient's previous value and/or a 15 normal level and/or a normal baseline level for the particular individual indicates therapy efficacy and/or decreased tumor burden.

The effect of any exogenous immunotherapy compound may depend upon its method of delivery which, in turn, may 20 determine how rapidly the compound is present at the site of action (sometimes termed target site). The target site may be one or more types of leukocytes that are classically regarded as involved with cell-mediated immunity, i.e., lymphocytes, e.g., T-lymphocytes, B-lymphocytes, or other cell types such 25 as macrophages, natural killer cells, mast cells, basophils, dendritic cells, etc. It will be appreciated that many if not all of these cell types may contain genetic modifications to affect their normal physiological function in the immunologic process. For example and without limitation, genetically modified enhancement, decrease, attenuation, facilitation of various targeting and activation function may be achieved using techniques known in the art. Because of the myriad types of administration routes coupled with the myriad types of exogenous immunotherapy compounds, delivery may be a consid-35 eration in patient therapy. The exogenous compounds may be delivered in a number of ways. One embodiment provides nanoparticle delivery, with nanoparticles described herein and as known in the art, and which include but are not limited to, quantum dots.

As described herein, nanoparticles may be biocompatible or rendered biocompatible. In one embodiment, nanoparticles may be associated with liposomes. Nanoparticles may contain and/or be coated with, in whole or in part, materials such as ferric oxide, carbon, diamond, zinc oxide, gold, etc. 45 providing methods to target the nanoparticles to a particular cellular, physiologic, and/or anatomic site. For example, one embodiment described herein uses an alternating magnet or electromagnetic radiation with a metal-containing nanoparticle (e.g., ferromagnetic nanoparticle) to effect therapy (see, 50 e.g., FIG. 5).

Nanoparticles are a particularly advantageous delivery vehicle for several reasons. Their small nanoscale size permits their entry, localization, and/or accumulation into anatomical and physiological regions otherwise less readily 55 accessible. Their nanoscale size likewise permits intravenous administration, providing systemic access via the circulatory system.

Permutations of any of these embodiments are included in the inventive method. Illustrative but non-limiting examples 60 include polymer-coated nanoparticles, nanoparticle-stabilized liposome-polymer nanocarriers, nanoparticles contained in or formulated with liposomes, micelles, reverse micelles, etc.

As described herein, the nanoparticles contain a targeting 65 agent, such as an antibody, a specific binding pair member, etc., ensuring the nanoparticles administered to a patient are

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directed to a target site, e.g., a tumor site, a particular cellular receptor, etc. Using this approach ensures the thermotherapy is targeted.

In one embodiment, a cancer patient receives immunosuppressive therapy. Such immunosuppressive therapy may involve chemotherapy, radiation therapy, or both chemotherapy and radiation therapy. Prior to the patient receiving immunotherapy, the patient provides a blood sample from which leukocytes are separated. The leukocytes are then cultured in vitro in an environment to support their growth and maintenance; T cells are treated with the cytokine interleukin-2 (IL-2). The cultured autologous leukocytes, termed lymphokine-activated killer cells (LAK cells), are then reinfused into the patient, providing a bolster to the patient's suppressed immune system. In this embodiment, the combination of hyperthermal therapy with immunotherapy may be used to advantage. This is because the patient receives combination therapy: (i) chemotherapy and/or radiation therapy, which may or may not be finely targeted to a specific malignancy, tumor type, tumor location, etc.; (ii) hyperthermal therapy to destroy cells at increased temperature that does not result in protein denaturation thus sparing normal cells and normal cellular processes, and provided at a specific location to which the therapy is directed at a target site; and (iii) immunotherapy to bolster or restore or augment the patient's own immune response and thus bolster or augment natural immune functions. The combination therapy avoids total immunosuppression and provides the patient at least a modicum of immune function during recovery. The combination therapy permits a decreased dose of nanoparticles administered, decreased frequency of hyperthermal therapy, or both, providing less of a therapy burden to the patient. The combination therapy, because it considers the immune system, facilitates patient recovery or comfort with fewer complications of the immunosuppressive treatment itself.

To effect therapy, one or more large energy generation sources or units can provide a source of thermal energy for a large part of the body. Alternatively, a simpler, e.g., smaller, energy generation source or unit, e.g., a hand-held unit, can provide a source of thermal energy to labeled nanoparticles when brought close to the desired tissue area. In one embodiment, electromagnetic radiation is the energy source, e.g., from ultraviolet radiation up to radiofrequency waves. In one embodiment, ultrasound energy is the energy source, e.g., an ultrasonic unit focusing ultrasound waves to the desired area of the body to heat the nanoparticles. In one embodiment, an alternating magnetic field is used to heat the magnetic nanoparticles and treat the tumor. An alternating magnetic field is readily created, e.g., by a simple iron rod surrounded by a coil of wire with alternating current.

The device providing the energy source can be a magnet, microwave unit, a radiofrequency (RF) probe, an ultrasound probe, etc. The device can be any size. The device may be stationary, hand held, or introduced into the body by invasive or minimally invasive methods, e.g., introduced by a catheter. The device can be brought close, adjacent, or proximate to the target site. If the device is an alternating magnetic field, the oscillation of the magnetic field can range from a few Hz to gigahertz with appropriate magnetic field force according to the distance from the target. In one embodiment, e.g., 0.000001 Tesla to 11 Tesla, with a distance of 0.1 cm to 15 cm, are used.

In one embodiment, non-magnetic, magnetic, or paramagnetic nanoparticles labeled with the tumor antibody and/or fluorescent moieties are used to selectively bind to tumor cells.

In one embodiment, the method is used to effect therapy of a non-malignant tumor. In one embodiment, the method is used on normal cells with a propensity or predisposition to malfunction and/or grow excessively. One such example includes myofibroblasts, which are activated after coronary stent placement. Another such example includes nerve cells, which may be aggravated and thus produce chronic pain. In these cases, a cell specific antibody is combined with nanoparticles and conjugated ACPPs to attach nanoparticles to axons of these nerves. The nerves are then treated non-invasively to achieve a localized temperature up to about 60° C. to effect selective ablation and, thus, to prevent transmission of pain sensations. Another such example uses the method therapeutically to stimulate tissue healing, e.g., a beneficial thermal effect in creating heat shock proteins and encouraging 15 circulation in the area.

In one embodiment, after injecting the nanoparticle composition, either with nanoparticles alone or combined with ACPP, the patient target(s) is exposed to an energy source (electromagnetic radiation energy, ultrasound energy, an 20 alternating magnetic field, etc. as previously described) to increase the temperature of the cells that are complexed with, i.e., that contain, the nanoparticles. In one embodiment, selection of a desired temperature, e.g., 39° C. to 58° C., duration, and/or imaging means as subsequently described, 25 selectively damages the tumor cells while protecting the normal cells. With surgical tumor excision, the tumor margins, which contain nanoparticles, are visualized by imaging means, e.g., photoacoustic imaging, magnetic resonance imaging, thermal imaging, etc. A magnetic resonance imaging unit can be used for imaging, alone, or for thermal imaging of the tissue. Alternatively, photoacoustic sound can be modified, e.g., amplified, so the sound can be heard by the surgeon during therapy and be used as a diagnostic test for the presence of the remaining tumor cells in the surrounding 35 unexcised tissue, guiding further treatment.

The method also permits remote treatment of a patient in need of such treatment but unable to travel for on-site treatment by electronic communication, e.g., Internet communication. The communication means is operatively linked to 40 software that controls an energy source and an imaging system. The system is integrated to provide an energy source as previously described, e.g., ultrasound, electromagnetic radiation, alternating magnetic field, and a thermal imaging system as previously described, e.g., photoacoustic or MRI. Using 45 this embodiment, medical personnel provide instructions that control both the imaging device and the heating device. When a desired temperature in the target tissue is achieved, the software modifies the thermal energy output, e.g., it maintains the tissue temperature at the desired thermal energy level or it 50 increases or decreases the thermal energy at its source. In one embodiment, the system is equipped with a tracking device and pattern recognition software to maintain an accurate location of the treatment site. In one embodiment, any variation, such as patient movement or a change in settings, triggers a 55 failsafe system to terminate the procedure. In one embodiment, the entire system provides real-time results to medical personnel at a location remote from the patient (e.g., down the hall or in a different country), and these medical personnel can control the entire process electronically by Internet con- 60 nection. The information is software controlled, medical personnel regulated, and subject to the above-described failsafe mechanisms and privacy standards as needed.

The target site is heated to the desired temperature to hyperthermally treat the target site using standard heating instruments and equipment for heating tissue and standard equipment for visualizing the dye in the target site that has been 42

released from the heat sensitive particles. For example, the heating equipment preferably includes suitable heat or energy source that is able to focus the heat or energy on the target and is able to control heat and temperature of the tissue. The heat source can be an electrical resistance heating element, or an indirectly heated element. The heating device can also have a radiation energy source for producing heat at the target site, such as a radio frequency (RF) device, ultrasonic generators, laser, or infrared device. One example of an RF generator device for hyperthermally treating tissue in a selected target site is disclosed in U.S. Pat. No. 6,197,022, which is hereby incorporated by reference in its entirety. Examples of suitable ultrasonic devices for delivering ultrasonic hyperthermia are disclosed in U.S. Pat. Nos. 4,620,546, 4,658,828 and 4,586, 512, the disclosures of which are hereby incorporated by reference in their entirety.

In one embodiment, the duration of the applied radiation energy may be from about a femtosecond to about 15 minutes. In one embodiment, the duration of the applied radiation energy may be from about one picosecond to about 15 minutes. In one embodiment, the duration of the applied radiation energy may be from about one nanosecond to about 15 minutes. In one embodiment, the duration of the applied radiation energy may be from about one microsecond to about 15 minutes. In one embodiment, the duration of the applied radiation energy may be from about 1 second to about 15 minutes. In one embodiment, the duration of the applied radiation energy may be from about 1 second to about 15 minutes. In one embodiment, the duration of the applied radiation energy may be from about 1 second to about 15 seconds.

In one embodiment, the temperature at which the radiation energy is applied ranges between about 45° C. to about 49° C. In one embodiment, the temperature at which the radiation energy is applied ranges between about 39° C. to about 58° C. In one embodiment, the temperature at which the radiation energy is applied ranges between about 39° C. to about 45° C. In one embodiment, the temperature at which the radiation energy is applied ranges between about 45° C. to about 49° C. In one embodiment, the temperature at which the radiation energy is applied ranges between about 49° C. to about 56° C. For example and without limitation, relatively brief treatment times are used for circulating cells (e.g., cells in vessels of the circulatory and lymphatic systems). Relatively longer laser pulses may also be used for tissues located deep inside the body. In one embodiment, the ultrasound frequency can range between 0.5 KHz to 200 MHz. In one embodiment, the ultrasound frequency can range between 0.5 M Hz to 10 MHz. In one embodiment, the ultrasound frequency can range between 10 MHz to 30 MHz. In one embodiment, the ultrasound frequency can range between 5 MHz to 80 MHz.

In one embodiment, the heat source includes a probe having a tip with the heating element or energy emitting element attached thereto. The energy emitting element can be an optical fiber operatively connected to a laser, infrared or ultraviolet light source. The probe preferably includes a suitable control mechanism for manipulating the probe to the target site and a control for controlling the energy applied to the target site. In one embodiment the wavelength of light is selected to be in the range between 350 nm to 1300 nm. In another embodiment the wavelength of light is selected to be in the range between 450 nm to 600 nm.

A suitable device for hyperthermally treating the tissue in a target site is shown in the figure. The device 10 includes a probe 12 having an optical fiber 14 with a distal end 16 for emitting a laser light to heat the tissue 17. Preferably, the end 16 of optical fiber 14 can focus the light source on the target site 17. Optical fiber 14 is connected to laser generator 18 that is able to generate a laser beam of sufficient intensity and

within wavelength for hyperthermally treating tissue. For use of the method in making a diagnostic assessment or for therapy, the tissue is treated to a temperature of at least 41° C. to 56°C, and preferably at least 42°C. to 56°C. In a preferred embodiment, probe 12 includes a second optical fiber 20 having a distal end 22 and a third optical fiber 24 having a distal end 26. Optical fiber 20 is operatively connected to a light source 28, such as a laser, that is able to emit a light beam having a wavelength capable of fluorescing a fluorescent dye in the target area when the dye is released from the heat 10 sensitive particles. Optical fiber 24 is operatively connected to a suitable imaging device 30 for capturing the fluoresced light from the excited dye and visualizing and producing an image of the fluorescing dye in the target site. Imaging device 30 can be a CCD or a device equivalent to a funduscope. An 15 example of a suitable funduscope is disclosed in U.S. Pat. No. 4,891,043 to Zeimer, which is hereby incorporated by reference in its entirety.

In another embodiment of the invention, the probe can include a heating element or a device for receiving a heated 20 fluid that can transfer the heat to the tissue in the target site. The probe can include an expandable bladder member for receiving a heated fluid delivered from a fluid-heating source. In still another embodiment, the expandable bladder includes a permeable portion so that the heated fluid can be applied 25 directly to the target site. A suitable aspirating device is preferably included to remove the excess heating fluid when applied directly to the target site.

In one embodiment, the target site is the retina or choroid in the eye of the patient. The heating and visualizing instrument 30 includes a laser capable of focusing a laser beam on the target site where the laser beam has a wavelength and intensity to heat the cells to a temperature of at least 42° C. In one embodiment, the laser heats the cells to a temperature of 50° C. or below and preferably to about 42° C. to 56° C. The 35 instrument also includes or is used in combination with a funduscope to excite or fluoresce the dye that has been released in the target site to capture and visualize the fluorescing dye. A funduscope that can be used is disclosed in U.S. Pat. No. 6,248,727, which is hereby incorporated by 40 reference in its entirety. The laser source is selected to provide sufficient energy to heat the tissue in the target site to the desired temperature.

The fluorescent dye is encapsulated in a suitable heat sensitive particle and introduced into the patient in a location to 45 be visualized in the target site. The heat sensitive particles can be microcapsules, or nanocapsules that are able to release the dye at a temperature of about 41° C., and preferably 42° C. or higher. In preferred embodiments, the fluorescent dyes are incorporated into heat sensitive liposomes that have a phase transition temperature at the temperature of hyperthermia. In one embodiment, the liposomes have a phase transition temperature within the desired temperature range that tissue or cells are to be heated.

In one embodiment, the liposomes have a phase transition  $^{55}$  temperature of at least  $41^{\circ}$  C. and preferably at least  $42^{\circ}$  C. In a preferred embodiment, the liposomes have a phase transition temperature of about  $45^{\circ}$  C. to about  $50^{\circ}$  C.

The liposomes can be made by various processes as known in the art. The phase transition temperature of the phospholipid is selected to control the temperature that the dye and other components are released from the liposomes. Phospholipids are known to have different phase transition temperatures and can be used to produce liposomes having release temperatures corresponding to the phase transition of the 65 phospholipids. Suitable phospholipids include, for example, dimyristoylphosphatidyl choline having a phase transition

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temperature of 23.9° C., palmitoylmyristoylphosphatidyl choline having a phase transition temperature of 27.2° C., myristolypalmitoylphosphatidyl choline having a phase transition temperature of 35.3° C., dipalmitoylphosphatidyl choline having a phase transition temperature of 41.4° C., stearoylpalmitoylphosphatidyl choline having a phase transition temperature of 44.0° C., palmitoylstearolyphosphatidyl choline having a phase transition of 47.4° C., and distearolyphosphatidyl choline having a phase transition temperature of 54.9° C. Another suitable phospholipid is a synthetic  $\rm C_{17}$  phosphatidyl choline from Aventi Inc. having a phase transition temperature of about 48° C.-49° C.

The phase transition temperature and the release temperature of the liposomes can be selected by combining the different phospholipids during the production of the liposomes according to the respective phase transition temperature. The phase transition of the resulting liposome membrane is generally proportional to the ratio by weight of the individual phospholipids. Thus, the composition of the phospholipids are selected based on the respective phase transition temperature so that the phase transition temperature of the liposome membrane will fall within the selected range. By adjusting the phase transition temperature of the liposome membrane to the selected range, the temperature at which the liposomes release the dyes and other components can be controlled during hyperthermia.

The liposomes in one embodiment of the invention are preferably prepared so that the liposome membrane has a phase transition temperature of at least 42° C., and preferably about 42° C. to about 50° C. In a preferred embodiment, the liposomes leak or rupture at a temperature of about 49° C. or less, and typically between about 45° C. and 49° C. In one embodiment, the phospholipids have saturated acyl groups. For example, glycerophospholipids can be used that have two acyl groups having 8 or more carbon atoms and where at least one of the acyl groups have at least 10 carbon atoms and typically 12-18 carbon atoms. Examples of suitable phospholipids include hydrogenated lecithin from plants and animals, such as egg yolk lecithin and soybean lecithin. The phospholipid can also be phosphatidyl choline produced from partial or complete synthesis containing mixed acyl groups of lauryl, myristoyl, palmitoyl and stearoyl.

The liposomes can be prepared from a mixture of dipalmitoylphosphatidyl choline and disteroylphosphatidyl choline in a weight ratio of 95:5 to about 5:95 and generally about 80:20 to about 20:80. In one embodiment, the liposomes are made from a mixture of dipalmitoylphosphatidyl choline and disteroylphosphatidyl choline in a ratio of 45:55 to about 55:45 provide a phase transition temperature of about 46° C. to about 49° C.

The liposomes of the invention can be prepared by standard processes as known in the art. The liposomes can be unilamellar or multilamellar and have a particle suitable for delivering the dye to the target site. In one embodiment, the liposomes have a particle size of a sufficiently small size to be introduced into the bloodstream of the patient in a location near the target site to flow through the target site.

The liposomes can contain a suitable osmotic pressure controlling agent that is physiologically acceptable to the patient. Examples include sodium chloride, sugars such as glucose, mannitol and sorbitol, and amino acids such as glycine, aspartic acid and glutamic acid. Examples of suitable process for preparing liposomes are disclosed in U.S. Pat. No. 4,235,871 to Papahadjopoulos et al. and U.S. Pat. No. 4,522, 803 to Lenk, which are hereby incorporated by reference in their entirety.

The liposomes of the invention contain a dye that is able to fluoresce and that can be visualized in the target site by exciting with a light source that is amenable to the target site and the patient. The fluorescent dye can be any fluorescent that is suitable for encapsulation and is physiologically compatible. Preferably, the fluorescent dye is quenched when encapsulated at an appropriate concentration. The quenching concentration is a sufficiently high concentration to mask or minimize detection of fluorescence when illuminated by a fluorescing light source. The quenching concentration can be 10 determined by routine experimentation as known in the art. When heated, the heat sensitive liposomes rupture or leak the dye and other components of the liposome so that the dye is diluted in the target site to a suitable concentration where the dye can be fluoresced and visualized upon excitation by a 15 suitable light source. Examples of suitable dyes include 6-carboxyfluorescein and its derivatives. Suitable fluorescent dyes can be excited by an emit light at wavelengths that are not strongly absorbed by the tissue. Other suitable dyes include indocvanin green and aluminum phthalocyaninetet- 20 rasulfonate. It will be understood that the fluorescing light source and the visualizing instrument are selected according to the wavelength of the fluorescing dye to visualize the dye.

In one embodiment, the dye is selected to fluoresce in the presence of a light from an argon laser, a helium-neon laser or 25 infrared laser. Preferably the dye is selected to be compatible with the exciting light or laser source to fluoresce when subjected to the light or laser beam. A suitable dye is sold under the tradename D-275 by Molecular Probes, Inc. and fluoresces green when exposed to light from an argon laser at 484 30 nm. A dye sold under the tradename D-1121 fluoresces orange when exposed to a long wavelength laser light at 560-574 nm. One preferred dye is an infrared excitable dye Dilc<sub>18</sub>(7), which fluoresces at a wavelength of 740-780 nm.

In one embodiment of the invention, a fluorescent dye is 35 encapsulated in liposomes having a phase transition temperature of 42° C. to 50° C., and preferably about 45° C. to 49° C. In another embodiment, the liposomes have a phase transition temperature to release the dye at a temperature of about 46° C. to about 49° C. The liposomes are injected into the blood- 40 stream of the patient in a location where the liposomes flow to the target site. In some embodiments, the liposomes can be introduced directly to the target site intravenously, subcutaneously or topically. A hyperthermal heat source and a dye exciting light source are applied to the target site. The hyper- 45 thermal heat source, which is preferably a laser light beam, is focused on the target site to heat the tissue and the cells to a temperature of at least 42° C. to hyperthermally treat the tissue and kill the cells. The hyperthermal heat source also heats the liposomes to a temperature at least equal to the phase 50 transition temperature to release the dye. The fluorescing light source excites the dye so that the fluorescing dye in the target site can be detected and visualized. By encapsulating the fluorescent dye in liposomes having a phase transition of at least 42° C., the detection of the fluorescing dye provides a 55 positive indication to the operator that the desired tissue temperature has been obtained that is necessary to hyperthermally treat the tissue. The phase transition temperature of the liposomes is selected according to desired minimum temperature that the tissue is to be heated. The hyperthermia 60 energy source is applied to the target site for a time sufficient to treat the tissue to the desired level. Generally, the tissue is heated to a temperature of at least 42° C. for 1-15 minutes and preferably 1-10 minutes.

In one preferred embodiment of the invention, the lipo- 65 somes contain a suitable drug or photosensitizing agent. The drugs preferably show a synergistic effect when combined

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with the hyperthermia treatment of the invention. The release of the drugs from the liposomes provide an improved targeting effect by releasing the drugs by the heat source in the target site. Suitable drugs include antitumor agents such as cisplatin, carboplatin, tetraplatin and iproplatin. Suitable anticancer drugs include adriamycin, mitomycin C, actinomycin, ansamitocin and its derivatives, bleomycin, Ara-C, daunomycin, metabolic antagonists such as 5-FU, methotrexate, isobutyl 5-fluoro-6-E-furfurylideneaminoxy-1,2,3,4,5,6 hexahydro-2,4-dioxopyrimidine-5-carboxylate. Other antitumor agents include melpharan, mitoxantrone and lymphokines. Antitumor therapy involving biologics, e.g., DNA, RNA, protein, siRNA, genes or portions thereof, etc., may be administered.

The amount of the particular drug entrapped in the liposomes are selected according to the desired therapeutic dose and the unit dose.

In one embodiment, the method administers a drug or drugs and/or a radiotracer, either liposome-contained or not, contained in nanoparticles. The nanoparticles are formulated, using methods known by one skilled in the art, to release the drug and/or radiotracer when a certain temperature is attained. As only one example, the nanoparticles can be conjugated with thermosensitive polymers that will disassociate and hence will release the drug and/or radiotracer upon reaching and exceeding a given temperature.

Such thermosensitive polymers and methods of formulating these thermosensitive polymers with or in nanoparticles containing chemotherapy drugs and/or a radiotracer are known in the art. Embodiments of the disclosed method use combinations of thermosensitive polymers with, e.g., known pH sensitive polymers, magnetically controlled release agents, magnetoliposome agents, thermosensitive liposomes entrapping iron oxide nanoparticles, etc., to achieve controlled drug release. The following non-limited examples are expressly incorporated by reference herein and generally describe such polymers and methods. Alavarez-Lorenzo et al. describe temperature-sensitive chitosan-poly(N-isopropylacrylamide) interpenetrated networks that have enhanced loading capacity and controlled release properties. Alexander et al. describe variable architecture polymers that are temperature- and pH-responsive, termed smart polymers, including synthetic copolymers and polymer derivatives. Hoare et al. describe nanoparticles encased in poly(N-isopropylacrylamide) (PNIPAM) that are heated by magnetic induction. Mornet et al. disclose poloxamers and poloxamines that are suitable vehicles acid (PMA) polymers that are modified with thiol groups (PMA<sub>SH</sub>) and are suitable vehicles for delivery of the nanoparticles and anticancer drug(s). In one embodiment, the nanoparticles are conjugated with radiological imaging compounds, e.g., gadolinium used in magnetic resonance imaging. In one embodiment, the nanoparticles are conjugated with metabolically active compounds, e.g., F-18FDG used to visualize tumors and metastatic lesions using positron emission tomography-computed tomography (PET-CT) scanning. It is known that heating cells to a temperature greater than 60° C. denatures its proteins and/or coagulates its contents, but due to the specificity of the antibody targeting, only the tumor cells are thermally treated; normal cells are unperturbed. Thus, these methods are used either singly or in combination to achieve desired effects of targeting tumor cells, and specifically tumor stem cells that are otherwise resistent to treatment and/or that elude detection.

The specific tissue site to be thermally treated, containing the complex of polymer-coated drug-containing nanoparticle and antibody complex, is thermally treated by any appropriate modality (e.g., electromagnetic radiation, ultrasonic

radiation, alternating magnetic field, etc.) to a first temperature at which the polymer and/or liposome releases drug. In one embodiment, this temperature is achieved relatively early in thermal treatment, e.g., from 37° C. to 43° C. After release, the nanoparticles are further thermally treated to achieve or 5 exceed a second temperature, e.g., from 43° C. to 45° C., 43° C. to 58° C., 45° C. to 50° C., 50° C. to 58° C., 58° C. to 60° C., to greater than 58° C., or to greater than 60° C. Without being bound by a single theory, the first thermal treatment primes the tumor cells and the tumor stem cells. When the 10 patient is receiving or has received chemotherapy, the priming results in the tumor cells' enhanced susceptibility to both the effect of chemotherapeutic drug and the damage from the rise of thermal energy inside the tumor mass. When the patient is receiving thermal therapy singly, or before chemo- 15 therapy, the priming results in the tumor cells' enhanced susceptibility to the damage from the rise of thermal energy inside the tumor mass, then are further damaged upon administration of high doses of chemotherapy, either locally administered to target to the tumor site. At the relatively high tem- 20 peratures at or exceeding 43° C., thermal damage is relatively selective in that it occurs primarily to cells with higher metabolisms, i.e., tumor cells. The method thus leaves nontumor cells unperturbed or minimally perturbed by the treat-

Examples of suitable photosensitive (photosensitizer) agents include aminolevulinic acid, porphyrin derivatives, porpurine derivatives, NPE-6, ATX-10, plant derived photosensitizers. Other synthetic sensitizers such as SNET2 and Lutex can be used. Preferably, the photosensitizers are used in 30 non-toxic amounts. In other embodiments, the liposome compositions can contain liposomes that encapsulate a hyperthermic potentiating agent such as perfluorooctyliodide, perfluoperfluorotripropylamine, rotributylamine, perfluorooctylbromide. Examples of liposome encapsulated 35 potentiators are disclosed in U.S. Pat. No. 5,149,319 to Unger, which is hereby incorporated by reference in its entirety. Other bioactive agents that can be delivered to the target site by encapsulating in liposomes include anti-inflammatory agents, antibiotics, antibacterial agents, antifungal 40 agents, anti-neoplastic agents and antiparasitic agents. Examples of anti-neoplastic agents include aclacinomycins, chromycins and olivomycins. In another embodiment of the invention, the liposome composition contains a mixture of liposomes having different phase transition temperature to 45 release the components at different temperatures. In one embodiment, the liposome composition contains liposomes encapsulating a first dye and having a phase transition temperature of 42° C. to about 45° C. and liposomes encapsulating a second dye and having a phase transition temperature of 50 about 50° C. or higher. In one embodiment, the second dye is encapsulated in liposome that release the dye at a temperature range of 50° C. to 60° C. Preferably, the second dye is able to fluoresce at different color than the first dye so that the dyes are distinguishable. In this embodiment, the liposome com- 55 position is delivered to the target and the target site is subjected to hyperthermia temperatures. As the tissue in the target site is heated to at least 42° C., the first liposomes rupture or release the first dye so that the first dye is visualized and detected in the target site. The detection of the first dye in 60 the target site enables the operator to monitor the temperature of the tissue in the target site and to indicate that a hyperthermal temperature has been attained in the tissue at the target site. During hyperthermia, it is difficult to determine and monitor the actual temperature of the tissue and care must be 65 taken to avoid overheating of the tissue and denaturization of the proteins. In preferred embodiments of the invention, the

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hyperthermal treatment does not exceed the protein denaturization temperature. In this embodiment, the second liposomes are selected to rupture or release the second dye at or slightly below the protein denaturization temperature. In this manner, the second dye is released and visualized to provide the operator with an indication that the tissue is heated to the protein denaturization temperature. The heat source is then adjusted by the operator to reduce the energy applied to the target site to avoid protein denaturization.

In another embodiment, the liposome composition can contain several liposomes that can leak or rupture at different temperatures to release the dyes at incremental temperatures as the temperature of the target site increases. In one embodiment, the liposomes can be selected to leak or rupture the dye at 2° C. intervals between about 42° C. and 50° C. The dyes for each liposome can be different to fluoresce a different color so that the different colors indicate a different temperature of the target site. In other embodiments of the invention, the tissue in the target site can be irradiated by beta radiation from strontium or iridium isotopes. Gamma radiation from P<sup>32</sup>, iodine-95, and palladium-90 can also be used. The radioactive isotopes can be delivered as small particles to the target site in combination with the hyperthermia treatment.

In another embodiment, the method combines thermal 25 therapy, either with or without chemotherapy, with localized internal ionizing radiation therapy to treat a patient in need of such treatment. The conditions of thermal therapy are described herein, and chemotherapy is also described herein and is known by those skilled in this art. The addition of localized internal ionization radiation therapy damage or destroy the endothelial cells lining the vessels proximate the tumor site, in effect starving the tumor of its nutrient supply source. This embodiment of the method administers a composition of targeted nanoparticles that containing an antitumor antibody, a radioactive isotope, and may also contain drug, to a patient. The patient receives an internal dose of radiation from the administered radioactive nanoparticles, i.e., internal radiation therapy, in contrast to external radiation therapy where the patient receives radiation from an external radiation source. This combination of localized thermotherapy and localized internal radiation therapy, and optionally localized chemotherapy, i.e., drug therapy, increases the likelihood of eradicating both existing tumor cells and tumor cell progenitors, as well as damaging endothelial cells in adjacent tumor-associated vessels. The method thus results in more robust therapy and increased chances of patient survival.

This embodiment relies on thermotherapy provided at the target site, and internal radiation therapy at the target site. In general, and as subsequently described, a composition of anti-tumor cell antibody associated with radioactive nanoparticles is administered to the patient, e.g., by intravenous injection. The tumor cell specificity of the antibody localizes the composition at the tumor site for thermotherapy, i.e., controlled hyperthermal therapy, and the associated radioactive nanoparticles damage or destroy the vessels adjacent the tumor, e.g., blood vessels, lymphatic vessels.

Thermotherapy is administered under conditions sufficient to result in targeted therapy to cells in the patient, as described herein. Radiation therapy results in damage to the tumor cells and adjacent endothelial cells in vessels providing blood, lymph, etc. to the tumor, i.e., to vessels feeding the tumor. As one example, nanoparticles can be coated with antitumor antibody to target, e.g., glioma cells in a patient with a brain tumor; an antibacterial antibody and an appropriate antibiotic, etc. Nanoparticles may be conjugated either directly with radioisotopes, or radioisotopes may be conjuaged with

polymers that are subsequently used coat the nanoparticles. Monoclonal antibodies may be used to attach coated nanoparticels to tumor cell surface markers, bacterial cell surface markers, etc. Nanoparticles are then provided, e.g., intravenously injected using a needle or catheter, applied through an 5 external orifice, to target the desired location. The dose can vary, e.g., from 10 μCi to 20,000 μCi. Radioisotopes producing a radiation are preferred, examples of which are At<sup>211</sup> (a particle, duration 7.2 hr, penetration 0.08);  $Ac^{225}$  ( $\alpha$ ,  $\beta$  particle, duration 10 days, penetration 10 0.1 mm);  $Bi^{212}$  ( $\alpha$ ,  $\beta$  10 particle, duration 60.6 min, penetration 0.09);  $Bi^{213}$  ( $\alpha$ ,  $\beta$ particle, duration 46 min, penetration >0.1 mm); Ra<sup>223</sup> ( $\alpha$ ,  $\beta$ particle, duration 11.4 days, penetration >0.1 mm);  $Pb^{212}$  ( $\alpha$ ,  $\bar{\beta}$  particle, duration 10.6 hr, penetration >0.1 mm); Tb<sup>149</sup> ( $\alpha$ , β particle, duration 4.2 hr, penetration >0.1 mm);  $I^{131}$  (α, λ 15 particle, duration 193 hr, penetration 2.0 mm);  $Cu^{64}$  ( $\alpha$ ,  $\lambda$ particle, duration 193 hr, penetration 2.0 mm);

 $I^{131}$  ( $\alpha$ ,  $\lambda$  particle, duration 193 hr, penetration 2.0 mm); and numerous other a emitters that can be similarly used, e.g., Bi<sup>213</sup>, B<sup>212</sup>, the boron neutron capture therapy (BNCT) that 20 can be used for superficial radiation penetration of tumor cells or adjacent cells to a depth of 7 µm. Similarly, groups of non-magenetic, paramagnetic, or magnetic nanoparticels can be produced. Polymeric coating of these nanoparticles renders them less toxic to normal tissue. Iodine<sup>131</sup> may be 25 crosslinked with antibody-coated nanoparticles to label the tumor or bacterial cells, both for therapy and imaging. Antiintegrins or anti-VEGFs may also be incorporated. The nanoparticles may be used simultaneously for drug delivery purposes against these tumor or bacterial cells, with the drug in 30 PDA, PGLA, dextran, dendrimers, PEG, etc. The nanoparticles may be quantum dots, polymer based nanoparticles, colloidal gold nanoparticles, iron oxide nanoparticles, etc. The nanoparticle size may range from 2 nm to 400 nm in one embodiment, and from 5 nm to 200 nm in another embodi- 35

Alpha-emitting isotopes emit radiation for a relatively short time, thus their effect will be short lasting and not damage normal tissue adjacent to a tumor. Isotopes of  $\alpha$  radiation emitted penetrate the tissue for a relatively short 40 distance. Because the nanoparticles are attracted to the tumor cell membranes, radiation affects not only the tumor cell but also the adjacent faxcular endothelial cells. The endothelial cells are specifically very sensitive to the radiation, therefore their obliteration deprives the tumor cells of their nutrient 45 source.

This treatment combination disrupts the vessels, to narrow the vessel lumen and/or damage the vessels themselves, killing both the tumor cells by the controlled thermotherapy with optional chemotherapy, and disrupting the nutrient source for 50 the tumor and progenitor stem cells.

In one use of this embodiment of combined thermotherapy and radiotherapy, vessels are effected such that the neovascularization in wet age related macular degeneration (wet AMD) is treated. To treat AMD, local choroidal injection is 55 used. The dose can vary depending on the mode of administration, body mass, body size, location of the tumor, etc. A does may range from 10  $\mu$ Ci to 20,000  $\mu$ Ci.

The radioactive nanoparticles for localized internal radiation therapy may be ferromagnetic or nonferromagnetic, as 60 described herein. They may be coated and/or otherwise conjugated, as described herein, with one or more anti-tumor cell antibodies, as described herein, then intravenously injected into or otherwise administered to the patient. The antibody conjugated and/or coated radioactive nanoparticles then 65 selectively bind to the targeted tumor cells, with the nanoparticles emitting either alpha radiation, gamma radiation, or

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beta radiation within a localized diameter of 1 mm to 2 mm from the site. The 1 mm to 2 mm radiation penetration range self-limits the radiation effect to the tumor-localized site, while desirably damaging endothelial cells adjacent the tumor. The radiation range damages endothelial cells in vessels feeding the tumor, diminishing or obliterating its nutrient source and in effect starving the tumor cells, as well as damaging tumor progenitor cells. The method targets all sites of tumor cells, and therefore treats both the primary lesion site and any metastatic sites.

Thermotherapy in combination with radiotherapy is performed using a controlled energy source, e.g., computer controlled, as described herein. A local magnetic and/or electromagnetic field is applied and monitored as described herein, e.g., MRI or photoacoustic imaging. In this way, the targeted cells and associated and proximate vasculature are affected. Non-targeted cells and distal vasculature, i.e., more than 1 mm to 2 mm from the tumor localized site, are not affected.

Another embodiment of the invention images the heat (temperature) production inside the eye (target) tissue. The desired temperature is achieved using a laser and photoacoustic imaging technique.

It is known that light can generate sound waves. This is the basis of photoacoustic technology. Some of the delivered energy, e.g., by laser, will be absorbed and converted into heat, leading to transient thermoelastic expansion and thus ultrasonic emission. The generated ultrasonic waves are then detected by ultrasonic transducers to form images. It is known that optical absorption is closely associated with physiologic properties, such as hemoglobin concentration and oxygen saturation. As a result, the magnitude of the ultrasonic emission, i.e., the photoacoustic signal, that is proportional to the local energy deposition, reveals physiological specific optical absorption contrast. Two- or three-dimensional images of the targeted tissues can then be formed.

A photoacoustic image is independently generated from an ultrasonic image, however, it is only visualized, i.e., imaged, by an ultrasonic receiver. Therefore, photoacoustic imaging and ultrasonic imaging should be performed simultaneously, so that the additional changes in the ultrasonic image, resulting from heat expansion, can be seen. In fact, if the ultrasonic wave and photoacoustic wave are parallel, there is no need for any other visualization system, as is the case is opaque tissue. The changes in the ultrasonic images are, however, dependent on the temperature that is created by an additional electromagnetic radiation, such as light, microwave generator, etc. This may be used when microwaves are used to treat intraocular tumors.

Presently, acoustically coupled resonant optical systems are sensitive enough to detect sound waves and differentiate the sound waves based on the temperature generated inside the target tissue. The pulsed light propagates in the ocular tissue uninterrupted until it meets the retino-choroidal tissue where it is differentially absorbed by the tissue producing a spatial distribution of the sound sources that can be imaged by an array of acoustic sensors. All visible light wavelengths and infrared wavelengths up to 1300 nm pass through the eye easily reaching the retina and choroids thus creating a high contrast between these tissue components. In the eye, most of retinal-choroidal pathology relates to the abnormal vessel formation, located at the junction of the retina and the choroids (new vessels), and most of the intraocular tumors are of retinal or choroidal origin permitting light, specifically infrared light at a wavelength of about 780 nm to about 1300 nm to penetrate these structures. The use of incoherent or partially coherent light permits penetration of a few centimeter and maintains a good spatial resolution for diagnosis or treatment.

The ocular tissue, from the cornea to the retina, provide a uniform optical density and index of refraction. This index of refraction changes when the light reaches the retina and choroids. However, the contrast image, in ultrasonic imaging, is related to the density and compressibility of the tissue, not the index of refraction, thus permitting photoacoustic imaging to be used in evaluating functional properties of certain molecules based on different optical absorption of molecules, e.g. in oxymetry differentiating oxygenated and reduced hemoglobin.

The diagnostic application of photoacoustic imaging is based on the absorption of electromagnetic energy by different molecules, producing different changes in temperature, pressure, and density. Therefore, photoacoustic image generation is the result of photothermal effect on the tissue or 15 molecules.

If the laser pulse is short enough, a local acoustic effect is generated that can be imaged by an ultrasonic transducer in 2D or 3D format. Because photoacoustic and ultrasonic imaging can share the same array and receiver, the image produced by them can simultaneously provide information on the thermal and anatomical structure, and location of the tissue in a rapid succession such as real time (video) images.

Nanosecond pulses can be generated from a Nd-YAG or Alexandrite laser. The laser delivery can be done either as a 25 combined transducer-laser head or independently through any optical system such as a slit lamp, a direct or indirect ophthalmoscope, or a fundus camera. These instruments have their independent illuminations permitting simultaneous visualization or imaging of the lesion in the eye using multiple imaging modalities, potentially along the previously described markers such as liposomes. In this case, the ultrasonic images are obtained through an independent transducer.

A contrast agent or a marker (biomarker) can be used to enhance the image or temperature (heat production) in the growing cells, such as choroidal neovascularization or tumor cells. For example, gold nanoparticles or tubes can be injected systemically which has a light absorption around 800 nm wavelength, which corresponds to a laser often used in ophthalmology for retinal coagulation purposes. The contrast generated by the above biomarkers in photoacoustic imaging is proportional to the concentration of the biomarker. The gold nanoparticles may be any shape, e.g., spherical, ellipsoidal, tubular (cylindrical). The gold nanoparticles may be solid or hollow. The size of the gold nanoparticles may range from 45 2 nm to 700 nm. In one embodiment, the size of the gold nanoparticles ranges from 50 nm to 250 nm.

In one embodiment, the thermal images, generated using photoacoustic imaging, can indicate progressively increasing tissue temperature while the area is being treated. In one 50 embodiment, a laser is used to treat the area while photoacoustic imaging is used to generate thermal images of the treatment area. In one embodiment, studies can be conducted to demonstrate the relationship between the photoacoustic images generated, as a result of a certain energy input, and 55 incremental temperature rise in the tissue to create a target temperature, for example, up to 55° C., or any other temperature below the temperature of protein denaturation.

Upon injection of labeled gold nanoparticles, their concentration in the peripheral blood increases. The concentration is 600 reduced within about 24 hours, depending in part on whether the gold nanoparticles contain (poly)ethylene glycol (PEG) groups or not (i.e., are PEGylated or not), because of hepatic clearance. The labeled gold nanoparticles are primarily either present in a bound form, e.g., an antibody-labeled-nanoparticle-cell (malignant cell) complex, or are absorbed by the tumor cells and hence are internalized. Hence, their decreased

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concentration in peripheral blood (e.g., blood sample obtained by venipuncture) indicates presence of a tumor cell, and provides quantitative information about the number of labeled cells per ml blood. Subsequently the total number of circulating cells in the total whole blood volume of a patient can be calculated. This permits discovery the presence of malignant cells, the ability to quantify the amount of circulating malignant cells from one lesion or multiple metastatic lesions which have metastasized, and to obtain information on their locations in the body.

Obtaining blood samples over a period of time (hours, days, weeks, etc.) and subsequently measuring the concentration of the labeled particles, indicates an increase or decrease in tumor cell shedding as a result of any therapy such as radiation, chemotherapy, thermotherapy, etc. It is thus of diagnostic usefulness, e.g., to monitor efficacy of therapy in a patient receiving therapy for a tumor. The patient receives an intravenous injection of a defined concentration of labeled gold nanoparticles. The concentration of the labeled nanoparticles in the peripheral blood is determined; this is a base level. Then, at a defined interval (e.g., days, weeks, etc.), as determined by a health professional, another peripheral blood concentration of the labeled nanoparticles is determined as an indicator of labeled nanoparticle clearance from the peripheral blood. The decrease in nanoparticle concentration is determined, and correlated with the previous uptake to determine the presence of a tumor, absence of a tumor, or decreased tumor burden (shedding). To illustrate, e.g., a decrease up to 30% to over 95% in 7 days indicates normal clearance of the labeled nanoparticles, and a decrease up to 99% over 7 days to 14 days indicates internalization of the labeled nanoparticles by a tumor. Any significant detection of nanoparticles beyond this time period indicates presence of circulating tumor cells; confirmation is obtained by evaluating a peripheral blood sample obtained by venipuncture and/ or by tumor cell histological examination.

In embodiments, the gold nanoparticles can be injected into a tumor, injected into a body cavity, applied over a mucosal surface, and/or applied to skin having a tumor. The absorption of the labeled particles can be used to differentiate a benign tumor from a malignant tumor and simultaneously to treat the tumor by the disclosed method of thermotherapy. In malignant tumors, the gold nanoparticles coated with antitumor antibody are absorbed by the tumor and remain in the tissue. In benign tumors, the gold nanoparticles coated with anti-tumor antibody may be initially absorbed by the tumor, but will not remain in the tissue and after 1-2 days are eliminated by systemic absorption.

In one embodiment, a method of delivering antibodycoated nanoparticles to a particular locus, or to a pathological site is disclosed. An example of a particular locus is the nervous system to treat a neurological pathology, e.g., a neurodegenerative disease. An example of a particular pathological site is a known tumor site, e.g., a breast tumor. Administration is either directly to or at the site (i.e., direct injection at the site) or indirectly (e.g., by injection into the bloodstream or other fluid surrounding a site, e.g., cerebrospinal fluid, ocular fluid, ventricles of the brain, etc.). Any site is amenable to the inventive method due to antibody directed targeting or localization. For tumor targeting, the method includes but is not limited to a superficial tumor site. Thus, the following examples are possible. The tumor may be located on an internal or external anatomical site, e.g., skin, extremities, breast, head, neck, and the like. The tumor may affect any organ, e.g., liver, kidney, brain, bone, prostate, ovary, tonsil, thymus, spleen, and the like. The tumor may be located in a deep tissue, such as in muscle, subcutaneous fat cells, intestines,

tendons, connective tissue, ligaments, intracranial, and the like. The tumor may be located at a site affecting body fluids, e.g., cerebrospinal fluid, ocular cavity, oral cavity, sinus or sinus cavity, lymph nodes, lymphatic vessel; or the site may be a bodily ventricle, e.g., heart ventricle, brain ventricle, and 5 the like; or the site may be a bodily cavity, e.g., bone cavity, uterus, kidney, bladder, pelvis, or the like.

In one embodiment, the antibody-coated nanoparticles are provided directly to the circulatory system, i.e., either injected into the arterial or venous blood vessel, or delivered by a catheter. In this embodiment, the method is used to treat pathological conditions of blood (e.g., erythrocytes, leukocytes, platelets, and/or their progenitor cells), and/or to provide systemic treatment, e.g., metastatic sites.

In one embodiment, the antibody-coated nanoparticles are 15 magnetic, diamagnetic, paramagnetic, and/or ferromagnetic.

In one embodiment, the method of providing antibodycoated nanoparticles to a site, or for systemic delivery, may be combined with immunologically based B-cells or T-cells that are optionally genetically modified. Such B-cells and T-cells 20 may attack localized and/or unlocalized cancer cells or hematological cancers (e.g., leukemias, lymphomas, etc.). In one embodiment, extra-corporal dialysis and/or plasmapheresis may be used in combination with the inventive method of delivering antibody-coated nanoparticles to accomplish 25 removal of excessive tumor protein in order to protect vital organs, e.g., kidney, liver, brain. In one embodiment, one or more of these therapies are employed at lower doses when used in combination with the method of delivering antibodycoated nanoparticles. In one embodiment, the method of 30 delivering antibody-coated nanoparticles may have a synergistic effect with another therapy, e.g., measured by metrics known to a person of ordinary skill in the art.

In other embodiments, nanoparticles of a material other than gold may be used. These include, without limitation, 35 diamond nanoparticles, platinum nanoparticles, combinations of gold, platinum, carbon, and/or diamond nanoparticles. Any of the above nanoparticles may contain at least one hydroxyl group. All such nanoparticles provide the various diagnostic and therapeutic applications as described above 40 for gold nanoparticles. The sizes and shapes are the same as those described for gold nanoparticles. All such nanoparticles may be covalently attached to (poly)ethyleneglycol, i.e., may be PEGylated.

All such nanoparticles create photoacoustic waves when 45 exposed to an external energy source such as light, ultrasound, lasers, radiation, microwave, etc. The temperature of the nanoparticles rises and their molecules expand. Molecular expansion produces an acoustic sound that can be recorded as a photoacoustic wave signal from an in vivo or in vitro envi- 50 ronment. These sounds, photoacoustic signals, are received by acoustic wave detectors or sensors, and are recorded and analyzed. Such methods are known in the art and are similar to signals obtained by endogenous chromophores such as hemoglobin. However, because the gold nanoparticles have 55 stronger absorption of the radiation than other chromophores, e.g., 30% to 99% higher, less energy is required to generate an appropriate signal. The signals of each chromophore can also be differentiated from other signals by ultrasonic spectroscopy. Acoustic wave sensors have acoustic wave resonator 60 elements including piezoelectric material (elements), as known by one skilled in the art. This ability to create and record the acoustic signal is useful for treating tumors, monitoring treatment efficacy, and making diagnoses, distinguishing malignant cells from benign cells, etc. in the same way as 65 previously described for optical tissue. For example, the absence of malignant cells does not generate an image in

photoacoustic imaging, and generates a different acoustic signature or characteristic in ultrasound spectroscopy. Thus, the method can evaluate the presence or absence of malignant cells in a human patient. An antitumor antibody-labeled nanoparticle, and a temperature indicating substance, in administered to the patient to form an antibody-labeled-nanoparticle-cell complex at the tumor site. Then, the tumor site is exposed to a radiation energy source under conditions sufficient to achieve a temperature of the complex between 41° C. to 56° C. The acoustic sound produced from the nanoparticle at the site is evaluated and correlated with the presence or absence of malignant cells at the target site by photoacoustic imaging and/or ultrasound spectroscopy.

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One embodiment using the above-described labeled nanoparticles is a method to differentiate a tumor or lesion containing malignant cells from one containing benign cells. The nanoparticles are coated with a targeting agent, e.g., antibodies other identifiers known to one skilled in the art, such that they target specific cells when intravenously injected or otherwise administered into a patient.

Cells that are labeled or tagged with gold nanoparticles show 30-40% increase in the resultant acoustic signal, compared to the untagged cells if the cells possess pigmentation. However, a sample of blood or other tissue, or a tumor (e.g., from biopsy) may be obtained and then bleached in vitro to rid the cell of pigment, e.g. melanoma cells. The samples are exposed to an external energy source (light, laser, etc) and the resulting photoacoustic wave signals are measured and analyzed using an ultrasonic spectrometer or other comparable device to determine the presence or absence of the malignant cells. The bleaching step is not needed for the majority of the neoplasms to generate photoacoustic signals, e.g., between a normal cell versus one tagged with gold nanoparticles. to be significant. The differences in the optical absorption of, e.g., light energy, permits differentiation, in vivo and in vitro, between normal and metastatic lesions (tumor cells).

In vitro acoustic cell analysis is performed by taking a biological sample from a patient, e.g., a known volume of blood or other body fluid that may contain circulating malignant cells, e.g., cerebrospinal fluid (CNS), lymphatic fluid, etc. To this fluid sample is added a specific antibody-coated gold nanoparticle. Because of the specificity of the antibody to a particular cellular receptor, protein, or other binding target, the gold nanoparticle binds to the specific cells present in the fluid and is detected, quantitated, monitored, etc. In one embodiment, an optional temperature-indicating substance is added for in vitro evaluation of the circulating tumor cells in the absence of nanoparticle injection. This is useful to indicate is metastatic cells are being released, because a positive result can initiate a search for potential metastatic lesions even if they have not progressed to a size so as to be visible by standard means of examination such as computed tomography (CT) or magnetic resonance imaging (MRI). The sensitivity threshold of recognizing a metastatic lesion is greater than about 1 mm diameter tumor, compared to a sensitivity threshold of greater than about 10 mm diameter tumor with standard means of examination. In either embodiment, the fluid is then analyzed by, e.g., an ultrasonic spectrophotometer or another appropriate device, to measure the number of tagged cells in the fluid. This procedure and analysis can also be adapted for use for other types of body sample such as tissue biopsies as known to one skilled in the art.

In one embodiment, a patient-specific antibody to a tumor is prepared. After biopsy, a specific antibody to the malignant cells of the tumor is generated. The antibody is then coated with gold nanoparticles resulting in an antitumor antibodylabeled nanoparticle. In one embodiment, a biphasic antibody

that binds a target via its Fab region is prepared. in one embodiment, an antibody that binds a target via its Fc region is prepared. Such procedures are known to one skilled in the art. These antibody-coated nanoparticles are specific for a specific tumor cell present in the patient forming an antitumor antibody-labeled nanoparticle-cell complex. In one embodiment, the complex may contain a compound, e.g., an antivascular endothelial growth factor and/or an antiproliferative agent. In one embodiment, the antibodies may also be labeled with a fluorescent dye.

As known to one skilled in the art, tumor cell membranes contain numerous receptors. In one embodiment, multiple antibodies against individual antigens may be generated from a tissue biopsy sample. Coating various gold nanoparticles with specific antibody against the tumor cell permits treating 15 a single tumor cells with multiple gold nanoparticles, either alone or with conventional chemotherapy and/or radiation therapy, thus enhancing the tumor-destroying potential of the method. In one embodiment, the combination of targeted hyperthermia plus radiation, etc. provides conditions in 20 which lower than typical levels of radiation are needed to destroy tumors, reducing radiation side-effects.

The above in vitro method and analysis can also be used to determine the temperature (energy) needed to kill tagged malignant cells, or other cells of interest, without damaging 25 the surrounding normal cells. This parameter is incorporated in an in vivo procedure to treat specific tumor or circulating malignant cells. For example, a tumor cell lacking pigmentation (e.g., breast cancer cells) may be killed using lower energy levels than a tumor cell having pigmentation (e.g., 30 melanoma cells).

In one embodiment, a collar positionable to fit on or around a patient's neck or extremities, contains a multiple diode laser that emits specific wavelengths of radiation energy to heat the labeled nanoparticles to a temperature sufficient to thermally 35 destroy the tumor cells and they circulate in the area encompassed by the collar. Other devices that can be used outside or implanted in the body as known to one skilled in the art may incorporate the method for treating and/or destroying malignant cells within other compromised areas of the body such as 40 those having tumors. Because the temperature generated inside the cells tagged with gold particles exceeds that of exposed non-tagged normal cells, this method protects normal cells while damaging the malignant cells.

In one embodiment, the energy source is a free electron 45 laser to provide the hyperthermal treatment. As described, the emitted energy from the free electron laser heats the nanoparticle of the described compositions to the desired temperature.

In one embodiment, the method uses magnetic energy to 50 provide the hyperthermal treatment. This embodiment is used for applying hyperthermal treatment to a lesion that is not accessible to application of light energy. Examples of such inaccessible lesions include brain cells, or cells deep inside the body.

In this embodiment, magnetic nanoparticles are used. In one embodiment, ferromagnetic, e.g., iron oxide or quantum dot, nanoparticles are used, either alone or combined with other magnetic materials or paramagnetic materials such as tungsten, cesium, aluminum, lithium, manganese, sodium, 60 platinum, and/or oxygen. Such magnetic nanoparticles are provided to a patient and exposed to an energy source that provides a magnetic field (e.g., coil, wire) that reverses the direction of the field rapidly using alternating current. In one embodiment, the particles are injected locally, e.g., inside a 65 tumor. In one embodiment, the particles are administered intravenously. In one embodiment, the particles are both

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injected locally and administered intravenously. In one embodiment, the described composition is introduced via an accessible cavity, such as oral, respiratory, or genitourinary, by injection or catheter. The particles heat up and expand, producing detectable acoustic (sound) waves. In one embodiment, the magnetic field ranges from about 0.0001 Tesla (T) to about 13 T. In one embodiment, the magnetic field ranges from about 0.0001 Tesla (T) to about 11 T. The degree of heat produced upon such exposure induce an expansion of the nanoparticles that is recorded by an ultrasonic transducer, as previously described.

In one embodiment, once delivered to the tumor site, or systemically injected or delivered, the particles are treated with an alternating magnetic field and/or electromagnetic radiation to heat or raise the temperature at the site. In one embodiment, the alternating magnetic field is from 0.0001 T to 13 T, or from 0.0001 T to 11 T, and the alternation frequency is from about 1 GHz to a few seconds, e.g., about 10 seconds, about 5 seconds, or about 2 seconds. In one embodiment, a temperature ranging from about 39° C. to 58° C. is achieved. In one embodiment, a temperature ranging from about 39° C. to about 45° C. is achieved. In one embodiment, a temperature ranging from about 42° C. to about 47° C. is achieved. In one embodiment, a temperature ranging from about 47° C. to about 49° C. is achieved. In one embodiment, a temperature from about 50° C. to about 58° C. is achieved. In one embodiment, the treatment is effected such that the increased temperature is maintained for a period of time ranging from about 5 seconds to 30 minutes. In one embodiment, the treatment is effected such that the increased temperature is maintained for a period of time ranging from about 1 minute to about 30 minutes. In one embodiment, longer times of maintenance at the increased temperature are used.

In one embodiment, simultaneous or substantially simultaneous imaging of the site, e.g., tumor, is effected by means known to the person of ordinary skill in the art, e.g., photoacoustic imaging, magnetic resonance imaging, X-ray imaging, optical coherence tomography, ultrasound imaging, fluorescence imaging, positron imaging, surface enhanced Raman spectroscopy (SERS), and the like. In one embodiment, the method of delivering antibody coated nanoparticles to a site is optionally combined with other therapeutic or medical techniques, e.g. a chemotherapy, radiation therapy, anti-vascular endothelial growth-factor therapy, steroid therapy, and the like.

The magnetic particles are attached to an antibody, as previously described, forming ferromagnetic-antitumor antibody-labeled nanoparticles. These particles provide diagnostic and/or therapeutic thermal heating at a biological target or lesion. A temperature indicating substance is also labeled with the antibody of interest, the ferromagnetic particle, or a combination of both.

As one example, a solution or formulation containing the ferromagnetic-antibody labeled nanoparticles is adminis55 tered by injection to a human or animal patient or by topical application. The antibody, upon binding to its cellular target, forms a ferromagnetic-antibody labeled nanoparticle-cell complex. After a period of time to allow clearance of the unbound ferromagnetic particles from the patient, the patient is placed in a magnetic field. In one embodiment, this clearance time is less than 2 hours. In one embodiment, this clearance time is 2 hours. In one embodiment, this clearance time is up to 48 hours. In one embodiment, this clearance time is up to 48 hours. In one embodiment, this clearance time is greater than 48 hours.

Clearance can be assessed the determining the presence of circulating nanoparticles in the patient. as follows. A small

volume of patient's blood is placed inside a small electric coil (0.0001 T-0.1 T) in vitro. Altering the magnetic field heats the particles and generates photoacoustic images measured by the ultrasonic transducer, which indicates the present of circulating nanoparticles. The pharmacokinetic of these circu- 5 lating particles can be measured. When the level of circulating nanoparticles are significantly reduced as described previously, e.g., 2 hours-24 hours post injection, the patient is positioned inside an a magnetic resonance imaging (MRI) instrument or any other applicable instrument known to one 10 skilled in the art.

The applied magnetic field heats the nanoparticles, including the sessile or attached particles to the tumor. The procedure is automatically controlled by photoacoustic imaging monitoring, as subsequently described below. The degree of heat production in the particles induces an expansion of the nanoparticles that is recorded by an ultrasonic transducer previously described. The patient is maintained in the magnetic field until a diagnostic or therapeutic temperature of 42° C. to 56° C. is reached. The time to reach this temperature 20 depends upon the energy applied in one session. The session can be repeated as desired. Generally, the exposure time may vary from <1 minute to >10 minutes. The acoustic sound produced from the nanoparticle at the site is evaluated and correlated with the presence or absence of malignant cells at 25 contains any gas. Because animal tissue always contains the target site by photoacoustic imaging and/or ultrasound spectroscopy. A photoacoustic image is created by exposing the tissue or site to electromagnetic radiation, e.g. radiation from 190 nm to about 10 µm; or microwave and radiofrequency radiation produced, e.g., using a laser. If the material 30 has magnetic properties, by reversal of the magnetic field, one can heat the tissue or site; one can also use mechanical waves produced by another ultrasound transducer to heat the tissue or site. These mechanisms create thermoelastic expansion of a molecule that generates a wideband (MHz) emission. The 35 time and amplitude of the photoacoustic images provide information regarding the absorption and location of the source, depending on the degree of thermal expansion of the material, etc., as known to one skilled in the art.

In one embodiment, the method uses magnetic energy to 40 provide the hyperthermal treatment. This embodiment is used for applying hyperthermal treatment to a lesion that is not accessible to application of light energy. Examples of such inaccessible lesions include brain cells, or cells deep inside the body.

In this embodiment, magnetic nanoparticles are used. In one embodiment, ferromagnetic, e.g., iron oxide, nanoparticles are used, either alone or combined with other magnetic or paramagnetic materials. In various embodiments, the described nanoparticle, e.g., ferromagnetic nanoparticle, is 50 carbon coated by methods known in the art, making the nanoparticle more biocompatible, e.g., less toxic. Such magnetic nanoparticles are provided to a patient and exposed to an energy source that provides a magnetic field (e.g., coil, wire) that reverses the direction of the field rapidly using alternating 55

The following methods control the energy applied to the nanoparticles, indicating when to treat after injection of the nanoparticles. A noncoagulative laser or radiofrequency or an acoustic wave, when absorbed, can produce a photoacoustic 60 ultrasonic emission of about 1 MHz or more (two- or threedimensional), creating an image of a tissue, site, etc. The image, in the form of a graph, curve, etc., shows an amplitude that depends on the amount of thermal energy absorbed or produced at the tissue or site. One can record this temperature 65 increase as an image, regardless of the exact degree increase. This characteristic can be used to distinguish low and high

energy applied, e.g. internally generated, by magnetic resonance on the ferromagnetic particles, and can be used to modify the amount of energy required. A computer processor (digital signal processor) can transmit the desired signal from the ultrasound unit to a device, e.g., a laser, radiofrequency device, magnet, etc., as known to one skilled in the art, to increase or decrease the applied energy to achieve potentially any desired temperature for any given situation and site. This energy varies; relatively low heat up to 55° C. for non-coagulative laser treatment, to >60° C. for coagulative laser treatment, to >100° C. in one embodiment and >500° C. in one embodiment for photoablative laser treatment, i.e., very short pulses but very high energy, depending on how deep the target tissue is located inside the body, and the property of the energy absorber inside that targeted tissue. This method results in a treatment that is visible, adjustable, and automatic depending on the needed temperature regardless of the depth of the lesion inside the body. Previously only a light source with an external photographic device could indicate the surface temperature. However these devices and wavelengths are limited by their wavelength, tissue penetration, and tissue water content, limiting light transmission to a few millime-

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Photoacoustic spectroscopy is very sensitive if a tissue gases as oxygen, nitrogen, and/or carbon dioxide, any degree of temperature increase causes expansion of these gases at that location. The sensitivity of photoacoustic spectroscopy can reach one part in trillion levels. Therefore this method demonstrates the thermal effect that was generated on the specific tissue, and the energy-absorbing nanoparticles can be imaged inside the body using this method.

As one example, a solution or formulation containing the ferromagnetic-antibody labeled nanoparticles is administered by injection to a human or animal patient human patient. The antibody, upon binding to its cellular target, forms a ferromagnetic-antibody labeled nanoparticle-cell complex. After a period of time, such as at least 2 hours up to 48 hours or more to allow clearance of the unbound ferromagnetic particles from the patient, the patient is placed in a magnetic field, such as that provided by an magnetic resonance imaging instrument or any other applicable instrument known to one skilled in the art. In various embodiments, the magnetic field is applied to a portion of the body, e.g., locally at the target 45 site, or to the whole body.

The applied magnetic field heats the nanoparticles. The degree of heat production in the particles induces an expansion of the nanoparticles that is recorded by an ultrasonic transducer previously described. The patient is maintained in the magnetic field until a diagnostic or therapeutic temperature of 42° C. to 56° C. is reached. The heat production is controlled to achieve this temperature and hence to treat only tumor cells. The acoustic sound produced from the nanoparticle at the site is evaluated and correlated with the presence or absence of malignant cells at the target site by photoacoustic imaging and/or ultrasound spectroscopy.

In embodiments, the described photoacoustic imaging may be combined with other imaging techniques such as X-ray, MRI, CT, and PET scans. In instances where the tumor is light accessible, optical coherence tomography (OCT) combined with photoacoustic imaging may be used.

In various embodiments, the method comprises providing a described nanoparticle composition, e.g., a nanoparticle labeled with an antibody, and providing an energy source which causes the nanoparticle to increase in temperature, resulting in heating tissue and cells in the tissue of an animal, particularly a human patient, at least to the temperature suf-

ficient to kill or damage the cells, in conjunction with photoacoustic imaging and/or ultrasound spectroscopy, where a desired temperature is achieved using an energy source and an imaging technique. As described, an energy source, such as a magnetic energy source, can be used which substantially penetrates the tissue, and thus allows activation of nanoparticle compositions, e.g., ferromagnetic nanoparticles, which may be located within deep tissue sites, or an energy source, such as light, may be used for surface or sub-surface activation of a nanoparticle composition. For example, in cases of surface tumors or tumors accessible by catheter, a carbon nanoparticle composition may be used which is activated or heated using light or electromagnetic radiation.

In one embodiment, any of the described methods for hyperthermally treating cells, including the use of magnetic 15 or gold nanoparticles, may be combined with platelet-derived therapy. In one embodiment, platelet-derived therapy comprises obtaining platelets from the patient's blood, coating the platelets with antibodies, and reintroducing the coated platelets into the patient. In one embodiment, the antibodies are 20 anti-tumor antibodies. In one embodiment, the coated platelets are reintroduced by injection into the patient's circulatory system after an initial hyperthermia therapy. The coated platelets travel though the blood vessels reaching the tumor surface where, due to the prior hyperthermia-induced tumor 25 cell damage, the injected coated platelets attach to the tumor and its associated vascular supply, creating a blood clot or thrombus, which obstructs the blood supply to the tumor, and nutritionally starves the tumor. In one embodiment, additional or subsequent hyperthermia therapy treatments may be 30 conducted during or following the platelet-derived therapy. In one embodiment, the platelets are coated with an antibody. This may be accomplished by methods as disclosed in, e.g., Iyer et al. (Meth. Mol. Biol. 2011; 751:553-563, Single-step conjugation of antibodies to quantum dots for labeling cell 35 surface receptors in mammalian cells, which is incorporated by reference in its entirety herein. Using such methods, cell surface receptors in platelets are labeled using antibody-conjugated semiconductor quantum dots. In one embodiment, the quantum dots are ferromagnetic, and as described above, 40 upon exposure to a magnetic energy source, provide hyperthermia therapy. The hyperthermia therapy provided by the platelet-coated antibody-quantum dot conjugate can be in addition to the hyperthermia therapy provided by the nanoparticle, or it can be the sole source for the hyperthermia 45 therapy. In one embodiment, any of the described methods and compositions, such as hyperthermia treatment with photoacoustic imaging, may be used for treating diseases or infections due to bacterial, fungal, viral, and/or protozoan organisms. In one embodiment, a described method may be 50 systemically applied, e.g., intravenous administration of the described composition. In one embodiment, the nanoparticle is labeled with an antibody which recognizes the infectious agent. In one embodiment, the nanoparticle is a ferromagnetic nanoparticle. In one embodiment, any of the described 55 nanoparticles are conjugated with an antibody that binds to, or associates with, an infectious agent, e.g., an anti-bacterial antibody, anti-viral antibody, anti-fungal antibody, anti-parasitic antibody, and the like. In this embodiment, the conjugated nanoparticle composition is locally or systemically 60 administered and subjected to radiation, such as electromagnetic radiation or alternating magnetic field, to achieve temperatures of between about 40° C. to about 49° C. in one embodiment, or between about 42° C. to about 49° C., or higher as needed to destroy the micro-organism(s). The toxins 65 from the infectious agent can be removed from the blood through a dialysis system. In one embodiment, the method

may be combined with additional therapies, including antibiotics, anti-fungal agents, anti-viral agents, etc. In various embodiments, following the method, the patient's blood is processed through an extracorporeal dialysis system, which removes or reduces the infectious agent, e.g., bacteria, and their associated toxins from the blood. In one embodiment, the dialysis system further comprises a magnetic separator, which allows separation and collection of nanoparticles, e.g., ferromagnetic nanoparticles, along with damaged infectious agent, e.g., bacteria, from the blood. In embodiments where a tumor is subjected to the described hyperthermia treatment, the circulating tumor cells can be collected by the magnetic separator along with the nanoparticles in the dialysis machine. In another embodiment, the patient's blood is exposed extracorporally to a localized alternating magnetic field prior to the blood being processed by a dialysis machine, such that the infectious agent, their associated toxins, and/or damaged tumor cells are separated and removed before recirculating the blood into the patient again, which prevents release of toxins inside the patient's body.

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In one embodiment, any of the described methods and compositions, such as hyperthermal treatment with photoacoustic imaging, are used to ameliorate a neurodegenerative disease, e.g., Alzheimer's disease, Parkinson's disease, and the like. In one embodiment, the described nanoparticle, such as colloidal gold or ferromagnetic, is conjugated with an antibody that binds to, or associates with, beta amyloid peptide which forms plaques found in the brain of Alzheimer's patients. In one embodiment, the neurodegenerative diseasetargeted nanoparticle is injected into the cerebrospinal fluid. The neurodegenerative disease-targeted nanoparticle may be subjected to electromagnetic radiation, such as microwave or RF, or alternating magnetic field, to achieve temperatures ranging between about 45° C. to about 55° C., for a period sufficient to reduce or prevent aggregation of beta amyloid peptides and/or to destroy beta amyloid peptide aggregates.

In one embodiment, the nanoparticle is colloidal gold. In various embodiments, the shape of the colloidal gold nanoparticle may be spherical, rod-shaped, irregular, etc. In one embodiment, the gold nanoparticle is rod-shaped, e.g., a nanorod, and may be subjected to surface plasmon resonance (SPR), which may be used as an imaging technique and/or as a means for heat generation for hyperthermia treatment, depending on the wavelength of the electromagnetic energy, e.g., near infrared light, and the axial diameter of the nanorod.

In one embodiment, the nanoparticle used in any of the described methods has (poly)ethylene glycol (PEG) groups attached. Methods to conjugate with PEG are known in the art and include, but are not limited to, encapsulating the nanoparticle and/or associating PEG with the nanoparticle. In embodiments, a targeting group is associated with the conjugated PEG; examples of such targeting groups include but are not limited to an antibody or a portion thereof, a ligand for a receptor, or other moieties that promotes association, retention, targeting, transport, and/or uptake of the described nanoparticle.

In one embodiment, the described nanoparticle, e.g., a pegylated colloidal gold or ferromagnetic nanoparticle, is configured as a drug carrier. In this embodiment, the nanoparticle is associated with a drug and facilitates or effects drug delivery by, e.g., drug administration, dispersion, targeting, controlled delivery (e.g., time-release), and the like. For example, hydrophobic drugs often require encapsulation or other means in order to increase the drug water solubility. In one embodiment, a drug is associated with a nanoparticle, such that the association may be covalent or non-covalent; in the case of a covalent association, the drug may be tethered to

the nanoparticle by a linking moiety. In one embodiment, a drug-associated nanoparticle is used in one of the described methods, such as hyperthermal treatment, in combination with a role as a drug delivery vector, such that, e.g., a target site (e.g., tumor) may be subjected to both hyperthermal 5 treatment and drug treatment,

In one embodiment, a thermoacoustic imaging technology is used with the tumor cells and tumor stem cells. Known photoacoustic technology uses light-induced heat, producing expansion, to create an ultrasonic sound that is recorded by an 10 ultrasonic transducer. The site is imaged as previously described. Photoacoustic technology is, however, limited by the penetration of the electromagnetic wavelengths from ultraviolet (UV) to far infrared (IR). The maximum penetration depth depends on the wavelength of the light that penetrates the tissue, e.g., 1-2 cm. Even with photoacoustic tomography (PAT), the maximum depth that can be imaged in the tissue is about 7-8 cm.

With the use of thermoacoustic imaging using heat sensitive antibody-coated nanoparticles and heating the coatednanoparticle complex (e.g., microwaves, radiofrequency waves, focused ultrasound, etc.) one can image the thermal expansion of intended tissue, e.g., tumor, etc. regardless of the tumor depth in the body. The same concept applied for creating an image using "photoacoustic"/thermoacoustic imaging, using magnetic antibody-coated nanoparticles with a reversible magnetic field. In both of these embodiments, MRI thermal imaging is also able to image the thermally treated tissue. Similarly one can heat the desired tissue with a focused ultrasound with one transducer, and observe the temperature 30 rise through the photoacoustic technology.

As described above, in various embodiments, a combination of hyperthermal treatment and imaging is provided. In one embodiment, the nanoparticle is colloidal gold which allows for hyperthermal treatment, as described above, as 35 well as imaging using SERS. In various embodiments, the colloidal gold nanoparticle is associated with quantum dots and/or Raman reporters. In one embodiment, the colloidal gold nanoparticle, associated with quantum dots and/or Raman reporters, is also associated with or encapsulated by 40 PEG. Further, as described above, the PEG may be further associated with a targeting moiety, such as an antibody or a portion thereof.

Thermotherapy Using a Magnetic Conformer

Magnets are found in nature and can be formed into any 45 shape. All magnets have the characteristics of absorbing other magnetic elements, e.g., iron. Electromagnets are produced by wrapping a wire as a coil around iron of any shape, size, or conformation. In an electromagnet the magnetic pole depends on the direction of the electrical flow in the surrounding wire and remains stable as long as electricity flows in the same direction. If the electrical flow is turned off, the electromagnet loses its magnetism. If the direction of the current is changed, the magnet changes its polarity and becomes an alternating magnet.

In one embodiment, the method uses a magnetic conformer. This conformer can be a natural magent or an electromagnet in which the magnetic field is not reversed. In such an embodiment, the treatment modality is applied either locally, systemically, or both locally and systemically 60 depending on the lesion and malignancy. When the treatment modality is applied systemically, the tumor has potentially metastasized in the body and the entire body is in need of thermotherapy. The localized magnet with stable magnetic field collects the circulating ferromagnetic nanoparticles or 65 nanowires in the tissue near its magnetic field. Nanowires, so-named because the width is smaller than the length, can be

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metallic (e.g., Ni, Au), semiconductor (Si, InP, GaN, etc.), organic, or inorganic and have greater efficiency converting light to energy.

In one embodiment, the localized magnet with a stable magnetic field attracts the ferromagnetic radioactive and/or drug coated nanoparticles or nanowires in tissue or in a benign or malignant tumor near its magnetic field. As in the general method disclosed herein, the nanoparticles or nanowires may be coated with anti-tumor antibody, contain therapeutic agent(s) (e.g., genetic agent (DNA, RNA), radioactive agent, chemical agent (chemotherapeutic, antimicrobial, etc.)) in any combination.

In one embodiment, the photoacoustic or thermoacoustic component of the thermotherapy system described herein is connected to a processor that is connected to the thermal energy producing system (FIG. 6). This embodiment permits specific control of the amount of energy delivered and maintained at the lesion site to create a specific temperature at the nanoparticle- or nanowire-cell complex site.

In one embodiment, the treatment modality is applied either locally or systemically but preferentially localized to a defined or partially defined tumor site with no systemic metastasis. The method is further disclosed using a choroidal melanoma of the eye, but as one skilled in the art will appreciate the method is not so limited and is applicable to practically any part and/or body system of the body (skin, mucosa, prostate, brain, vertebra, extremities, urogenital system, endocrine system, gastrointestinal system, pancreas, breast, etc.).

The general use of radioactive nanoparticles in a method for treatment of eye tumors has been fully disclosed previously. In the embodiment currently described, magnetic nanoparticles or nanowires and a conforming magnet, also termed a magnetic conformer, are used in conjunction with photoacoustically controlled temperature measurement. This is achieved by connecting the electromagnetic conformer, in this case as the source of energy delivery, to a processor that is in turn connected to the photoacoustic system, to adjust the desired temperature level at the lesion site and is to a predetermined level. As previously disclosed, energy sources can be electromagnetic radiation, microwave radiation, radiofrequency (RF) radiation, ultrasound radiation, an alternating magnet, etc.

In one embodiment using an electromagnet, a simple magnet is positioned externally at or near a tumor site, or internally at or near a tumor site by a minimally invasive surgical procedure to attract circulating ferromagnetic nanoparticles or nanowires. The magnet size depends on the tumor or lesion size. For positioning in the eye, the magnet will be relatively small in comparison for positioning in or near breast, bone, or soft tissues. The magnet shape, like magnet size, is variable and flat, concave, semicircular, circular, annular etc. shapes may be used depending on the tumor or lesion location.

The simple magnet is an electric coil that is positioned around a metal, with a positive pole and a negative pole. To produce an alternating magnet, an electrical switch is used to rapidly change the direction of electricity and change the magnetic pole of the metal with an alternating frequency of 1 per second to 1 per nanosecond and producing a magnetic field of, e.g., about 1×10<sup>-6</sup> Tesla to 10 Tesla. Prior to converting the magnet to an alternating magnet, the nanoparticles circulating in the blood or lymphatic fluid or provided at a specific site are concentrated in the area of the localized magnet with a stable magnetic field.

Using the electrical switch, the magnet is made to rapidly alternate its polarity, i.e., producing an alternating magnetic field. The nanoparticles or nanowires located in the alternat-

ing magnetic field of the coil develop a semicircular motion. The rapid polarization alternation creates heat in the nanoparticle or nanowire. Because the nanoparticles or nanowires are concentrated at or near the tumor site, the heat that is generated heats the nanoparticle-cell or nanowire-cell complex at or near the target site.

In one embodiment, the nanoparticles are injected into the circulatory system. While other administration methods are within the inventive scope, the ease and convenience of direct intravenous injection, or less routinely intraarterial, injection 10 makes this route attractive and provides a rapid route to reach all organs. Topical, intrathecal, subcutaneous, submucosal, inhalation, oral, cerebrospinal fluid injection, or intracavity (bladder, uterus, etc.) administration routes are possible to directly reach tumors or lymph nodes.

Regardless of the route of administration e.g. intravenously, of the nanoparticles, once in the body they are attracted selectively more to a magnet that is positioned in the localized area. Using the eye as a non-limiting example only, the conforming magnet can be placed externally on, or provided on a small probe to, an accessible ocular lesion, or the conforming magnet may be implanted on an internal lesion, e.g., under the conjunctiva, over the sclera, etc.

After a period of time post administration of 1 min to 15 min, sufficient to concentrate the nanoparticles or nanowires 25 at a target site, they are preferentially treated by the disclosed method of thermotherapy. The presence of the magnetic conformer converts the initially stable magnet to a reversible magnetic capable of generating localized heat in the nanoparticle-cell or nanowire-cell complex, without displacing their 30 location. In alternative embodiments, or after removing the localized magnet, the lesion or whole body undergoes hyperthermal treatment generated by means other than a conforming magnet, e.g., electromagnetic radiation, microwave radiation, radiofrequency radiation, etc., with the controlled 35 temperature using the photoacoustic processor system as previously described herein in detail.

In either embodiment, hyperthermal treatment involves generating heat from 37° C. degree to <60° C. in stepwise 1° C. to 3° C. increments gradually as desired for release of 40 agents from the nanoparticle and damage to the tumor cells. The polymers carrying the agents release them during the heating process, i.e., when the temperature for the polymer stretching or melting occurs sufficient to release the agent. In embodiments where a combination of treatment modalities 45 are applied, the combination of agent therapy and radiotherapy with thermal therapy provides synergistic treatment for the lesion or tumor.

The duration of therapy can vary from 0.5 minute to 1-2 hours, depending upon the duration and therapy needed, as 50 well as patient parameters, as one skilled in the art will be able to determine. In tumors that have metastasized, thermal treatment may be applied to the whole body after localized therapy. Using again the illustrative and non-limiting example of use of the method to treat an ocular lesion, the 55 conformer can be directly applied on the cornea or the conjunctiva when a lesion is externally located. To treat ocular lesions that are not externally located, e.g., tumors located in the back of the eye or close to the optic nerve, a conjunctival incision is used to provide the conforming magnet to the area. 60 The incision size depends upon the lesion size, and the magnet can be precisely located on or near the lesion site using, e.g., ophthalmoscopy, ultrasound, MRI, CT scan, PET scan, etc., as known to one skilled in the art. Once the location is verified, the magnetic conformer is temporarily placed and 65 stabilized, e.g., sutured in place, to the scleral surface over or in the vicinity of the lesion for a desired time sufficient to

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accumulate an adequate amount of nanoparticles or nanowires at that site. The treatment can be repeated at any time as desired, and gauged by use of one or several imaging methods described.

Exsosomes are univesicular or multivesicular structures originating from the plasma membrane of all cells, including cancer cells. Exosomes serve in cellular waste management, but can also function between adjacent cells in intracellular signaling, and as a conduit to transfer genetic material between cells. In some circumstances, exosomes can "infect" adjacent cells or travel in the circulation and facilitate a metastatic process. Booth et al., J. Cell Biol. 172 (2006) 923.

Exosomes are much smaller than the cells from which they originate, typically exosomes are between 30 nm-100 nm. Their membrane, similar to those of other cells, is composed of protein and lipid layers. Exosomes carry proteins and genetic components (e.g., RNA, DNA) and have the same surface receptors as the cells from which they originate.

Exosomes thus carry biomarkers of their cell of origin, either a normal cell or a cancer cell, and thus can serve as a marker for the existence of cancerous cells. However, exosomes also may be used to evaluate the prognosis of a pathology, e.g., malignant tumors, infectious organisms, viral infection diseases, etc. An increase of exosomes originating from cancerous cells in the vascular system can be an indicator that a cancer is more of an aggressive type. The presence of exosomes in the vasculature may facilitate metastasis. There is no known therapy to target such exosomes, either circulating or localized.

An embodiment of the invention is a method using hyperthermal therapy applied to target either circulating or localized exosomes. This embodiment of the method simultaneously recognizes, quantifies, and destroys or reduces the number of exosomes, using hyperthermal therapy. The method is useful for both diagnostic and therapeutic applications. Quantification of exosomes is used for diagnosis of, e.g., a malignancy, an infection, etc., and can be performed either in vitro and/or in vivo. Destruction or reduction of exosomes is accompanied by destruction or reduction of the cancer cells from which they are produced, and is used in therapeutic applications.

In cancer immune therapy, the patient's circulating tumor exosomes can be cultivated, using tissue culture methods known in the art, with a population of the patient's growing defensive cells: antigen-presenting cells (APC), dendritic cells, and/or macrophages in order to sensitize these defensive cells. Upon intravenous injection back into the patient, these now sensitized defensive cells activate the patient's own T cells to attack both the tumor cells and the circulating exosomes originating from the tumor cells. The disturbed microenvironment of the tumor after thermotherapy using labeled nanoparticles and/or quantum dots contribute to making the remaining tumor cells accessible to the sensitized T cells. In addition, RNA and DNA is released from the damaged exosomes in the serum and, in turn, is damaged by oxidative molecules such as hydroxyl radicals and other reactive oxygen species in the serum, important for treatment of viral and infectious diseases.

The nanoparticles and/or quantum dots may contain polymers such as (poly)ethylene glycol, i.e., they may be PEGylated, to increase their longevity and biocompatibility, as subsequently described. The nanoparticles and/or quantum dots may be associated with cell penetrating peptides to enhance their uptake in the tumor cells. As only one nonlimiting example, the nanoparticles and/or quantum dots may be associated and/or coated with a thermosensitive polymer (e.g., chitosan, chitin, etc.) to deliver compounds that are

anti-tumor or anti-cancer agents. Such compounds include, but are not limited to, antineoplastic agents. Such agents include drugs and/or biologics. In one embodiment, the nanoparticles and/or quantum dots can be conjugated with humanized antitumor monoclonal antibodies designed for the treatment of cancers and for the treatment of bacterial, viral, and parasitic infections.

The content of exosomes can influence the function of the cells in which they come into contact, or in which they are taken up. A plurality of organic, non-organic, or synthetic 10 nanoparticles alone or, with a plurality of quantum dots, are associated with various compounds to enhance their biocompatibility. Such compounds are known in the art and include, but are not limited to, (poly)ethylene glycol (PEG) and various PEG moieties, organic molecules, proteins such as biotin, 15 short peptides, naturally occurring amino acids such as arginine or phenylalanine, mono- or bilayers of phospholipids, or biotargeted molecules etc. Association includes coatings, covalent attachments, and other types of associations sufficient to enhance biocompatibility. For example, the nanopar- 20 ticles may be PEGylated to increase their longevity. The nanoparticles can consist of both inorganic or organic materials, as previously described. The particles can be entirely or partially biodegradable. The particles may also be included in or coated on a bioabsorbable or non-bioabsorbable but bio- 25 compatible polymer.

The nanoparticles are also conjugated with antibodies against the cell membrane's multiple exosome receptors and tumor cell membrane receptors. The conjugated and biocompatible nanoparticles, termed labeled nanoparticles, are 30 attached to and/or taken up by both exosomes and the malignant or normal cells.

In one embodiment, the inventive method may be used to damage the exosomes and count the circulating cells and exosomes in the blood with a wrist band or a neck collar 35 counter. Using the inventive method, one can also kill cells or damage exosomes simultaneously.

The labeled nanoparticles or quantum dots can be magnetic, paramagnetic, or nonmagnetic; gold, quartz, silicon, graphene, zinc oxide, ceramic, plastic, etc. Particles of crys- 40 talline silicon may be monocrystalline cells, poly or multicrystalline cells, or ribbon silicon having a multicrystalline structure. The particles may be a nanocrystal of synthetic silicon, gallium/arsenide, cadmium/selenium, copper/indium/gallium/selenide, zinc sulfide, indium/gallium/phos- 45 phide, gallium arsenide, indium/gallium nitride, and are synthesized controlling crystal conformations and sizes. In one embodiment, the nanoparticle may comprise a nanocrystal, such as cadmium/selenium (Cd/Se), and a metal. For example, a CdSe/Au nanometer-sized composite particle 50 may be synthesized through a two-step procedure, where CdSe nanorods are formed by the reaction of Cd and Se precursors in a mixture of trioctylphosphine oxide and an alkylphosphonic acid to form rod-shaped CdSe nanoparticles, and the CdSe rods are treated with a mixture of gold 55 chloride, didodecyldimethyl-ammonium bromide, and hexadecylamine to stabilize the nanocrystals and to reduce the gold chloride to elemental gold. Because the two ends of the CdSe rods differ crystallographically, and therefore chemically, control of growth conditions allows growth of Au par- 60 ticles preferentially on one end of each rod. In addition to CdSe/Au particles, one skilled in the art will readily recognize that particles can be constructed from a variety of other semiconductor/metal and semiconductor/semiconductor hetero-junctions. For example, particles based upon semicon- 65 ductor/metal hetero-junctions between group II-VI, IV, III-V, IV-VI, referring to groups of the periodic table, metal-oxide,

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or organic semiconductors and a metal, and in particular those based upon Si/Au, GaAs/Au, InAs/Au, and PbS/Au heterojunctions, can be used in the same way as those discussed here and one or more combinations of these can be used.

The size of the nanoparticles is between 1 nm-100 nm. In one preferable embodiment, the size of the nanoparticles is between 1 nm-10 nm. In this embodiment, the nanoparticle size is about an order of magnitude smaller that the exosome size.

Exosomes circulate in body fluids and, because of their small size, they cross the blood brain or blood ocular barrier and are taken up by normal cells or tumor cells. The labeled nanoparticles attach to the exosomes.

The labeled nanoparticles or quantum dots, when injected into the body, seek the exosomes and cell membrane receptors on cells such as tumor cells. As a result, the nanoparticle or quantum dot-conjugated exosomes are present in the circulation. The nanoparticles and/or quantum dots are excreted completely in the urine. As a result, the nanoparticle and/or quantum dot-conjugated exosomes can be detected extra corporally in a body fluid sample, e.g., blood, extra and intraocular fluid, cerebrospinal fluid, sputum, nasal secretions, urine, sweat, etc.

The complex of exosomes and nanoparticles and/or quantum dots can be detected and quantitated using various methods.

One quantitation and detection method uses that disclosed in U.S. Pat. No. 8,119,165 for detecting circulating malignant cells. The complex of nanoparticles and/or quantum dotsexosomes are heated using external thermal energy, e.g., electromagnetic, ultraviolet, visible, infrared, microwave, radiofrequency radiation, ultrasound, reversible magnetic field, etc. or a combination of these. The increased temperature in these complexes is measured using photoacoustic technology as described in U.S. Pat. Nos. 8,121,663 and 8,554,296. The thermal energy is preferentially absorbed by the nanoparticles and/or quantum dots, thus the temperature of the nanoparticles and/or quantum dots rises much more rapidly than the temperature of the tissues or cells to which they are attached or taken up. The temperature rise causes physical expansion of the nanoparticles and/or quantum dots, producing a sound wave that can be measured, imaged, and counted by a photoacoustic system. The temperature variation can also be measured precisely with this technique, demonstrating either an increase or decrease of the sound wave amplitude in photoacoustic imaging or tomography. The acoustic signal is measured and related to the temperature of the nanoparticle complex to effect hyperthermal therapy. The method can be combined with flow cytometry for cell quantitation. The quantitation and detection method can be used ex vivo, e.g., on body fluid samples.

Another ex vivo quantitation and detection method is by application of various wavelengths of light to the samples. The complex produces a visible response based on light exposure, e.g., fluorescence, and the visible response can be quantified by a detector and processed by a computer.

Other ex vivo quantitation and detection methods include standard methods that include flow cytometry, microscopy, mass spectroscopy, bioformatic analysis, etc.

In one embodiment, the nanoparticle and/or quantum dot complex may contain or be associated with various therapeutic compounds using a thermosensitive coating, e.g., a thermosensitive polymer as known in the art. Therapeutic compounds include, but are not limited to, anti-infective agents, anti-viral agents, anti-parasitic agents, anti-fungal agents, anti-proliferative agents, enzyme inhibitors or modulators, anti-VEGF, gene suppressants, gene stimulators, etc., as

known in the art. In this embodiment, increasing the temperature of the complex by hyperthermal therapy of nanoparticles and/or quantum dots and exosomes to about 39° C.-41° C. releases the compound from the thermo sensitive polymers associated with the nanoparticle/quantum dot carriers. Fur- 5 ther temperature increases to 43° C.-45° C. or 47° C.-50° C. or higher adversely affect the exosome membrane, rendering the exosome more vulnerable to the therapeutic agents released from the nanoparticles or administered at lower dose orally or systemically. This controlled thermotherapy can be combined with other standard therapies for cancer, e.g., radiation, immune, or vaccine therapy etc. Simultaneously a second plurality of nanoparticles and/or quantum dots may be administered containing other agents, e.g., small molecules, other therapeutics including but not limited to gene therapy, to 15 fight the disease process.

The temperature increase is related to the amount of energy applied. The photoacoustic system measures the temperature of the nanoparticle and/or quantum dot, and communicates by a processor with the energy delivery system to control the 20 amount of energy delivery to the complex of nanoparticles and/or quantum dots and exosomes at a predetermined level.

In one embodiment, the nanoparticles and/or quantum dots provides therapeutic agents in addition to its hyperthermal effect. In this embodiment, the nanoparticles and/or quantum 25 method by, e.g., radiofrequency, microwave, alternating magdots may provide immune-stimulatory agents, vaccines, and other therapeutic compounds known in the art. As only one non-limiting example, the nanoparticles and/or quantum dots may be associated and/or coated with a thermosensitive polymer (e.g., chitosan, chitin, etc.) to deliver compounds that are 30 anti-tumor or anti-cancer agents. Such compounds include, but are not limited to, antineoplastic agents. Such agents include drugs and/or biologics. Antineoplastic drugs include cisplatin, carboplatin, tetraplatin, iproplatin, adriamycin, mitomycin C, actinomycin, ansamitocin and its derivatives, 35 bleomycin, Ara-C, daunomycin, metabolic antagonists such as 5-FU, methotrexate, isobutyl 5-fluoro-6-E-furfurylideneamino-xy-1,2,3,4,5,6 hexahydro-2,4-dioxopyrimidine-5carboxylate, melpharan, mitoxantrone, lymphokines, aclacinomycins, chromycins, olivomycins, etc. Antineoplastic 40 biologics include DNA, RNA, protein, siRNA, genes or portions thereof, etc.

In this embodiment the immune response is stimulated, e.g., to facilitate removal of the hyperthermally-damaged tumor cells and their circulating exosomes, to target infec- 45 tious agents, etc.

Exosomes play a role in Alzheimer's disease by facilitating aggregation and accumulation of beta-amyloids proteins. Therapies that facilitate their removal would facilitate treatment. Stimulated microglial cells may be activated by adding 50 therapeutic agents to the complex of nanoparticles and/or quantum dots and exosomes. The complex may be administered orally, systemically, or injected into a body cavity. The use of combined nanoparticles and/or quantum dots and exosomes enhances cellular uptake of genes that can provide 55 combined genetic therapy with more conventional drug therapy, enhancing attach of a disease process.

In one embodiment, neurodegenerative diseases such as Alzheimer's disease or Parkinson's disease may be treated by the inventive method using nanoparticles and/or quantum 60 dots containing antibodies against beta-amyloid protein, e.g. Crenezumab, etc. These nanoparticles and/or quantum dots, which may be 1 nm-5 nm, pass through the blood brain barrier and penetrate into areas not previously accessible to such treatment. This embodiment of the method can be used for 65 treatment of degenerative disease of the central nervous system and/or the eye.

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In one embodiment, the nanoparticles are coated or otherwise associated with an antibody to specific cells, e.g., antitumor antibody, and are administered to a patient in need of therapy. The administered nanoparticles accumulate at a tissue site that is targeted by the antibody through typical antigen-antibody binding. The nanoparticles accumulated at the target site are radiated with an energy source, inducing a photoacoustic signal or sound wave from the nanoparticles. Using a processor, the photoacoustic technology controls the amount of thermal energy delivered at the desired temperature to the target site. The target site tissue is imaged, the temperature is recorded, and the photoacoustic sound wave is recorded, either from one location external to the body or from multiple locations. The recorded sound waves are amplified and processed to generate a computational tomographic image of the nanoparticles at the tissue target site. Since the antibody-coated nanoparticles are directed to attach to one or more surface receptors of specific cells, if the nanoparticles are attached to tumor cells, one obtains a two- or three-dimensional photoacoustic image of a tumor. If the antibodies are directed to normal cells of an organ, one obtains a two- or three-dimensional photoacoustic image of that organ regardless of its location.

In one embodiment, internal organs are imaged using the net, or focused ultrasound which penetrates deep in tissue to controllably heat the nanoparticles to obtain high-resolution two- or three-dimensional photoacoustic images of either a tumor or an organ.

In one embodiment, the temperature from the photoacoustic source is communicated to the energy source to control, via the processor, the amount of the energy and its duration as delivered to a desired degree of temperature measured at the nanoparticle tissue complex.

The thermal energy that is applied to the desired tissue target site, containing the accumulated nanoparticles, induces a photoacoustic sound. The photoacoustic sound is recorded and produced as each of a thermal graph of the target site, and as an image in any of one-, two-, or three-dimensions.

In one embodiment, a combination of imaging systems may be used. Illustrative and non-limiting examples include photoacoustic temperature imaging and ultrasound or focused ultrasound, photoacoustic temperature imaging combined with MRI or functional MRI (fMRI), photoacoustic temperature imaging combined by CT, PET, etc. scan, photoacoustic temperature imaging combined with photography or OCT, photoacoustic temperature imaging combined with alternating magnetic field imaging, etc. Because other imaging systems such as MRI, CT scan, PET scan etc. do not provide a high resolution image, their combination with photoacoustic imaging results in enhanced image acquisition with improved resolution.

The thermal energy applied may be in the form of electromagnetic radiation, ultraviolet radiation, visible light, infrared light, radiofrequency waves, microwaves, focused ultrasound, alternating magnetic field radiation, or a combinations of these. The combined use of various energy delivery systems decreases the amount of energy required for each unit or source, which may be desirable. Examples of combined therapy systems include, but are not limited to, lasers of various wavelengths, radiofrequency waves, microwaves with focused ultrasound, microwaves with alternating magnets, etc.

In one embodiment, the thermal energy may be applied from multiple sites to prevent pain to the patient and possible overheating of the target site. A single energy source can exceed the thermal pain tolerance when applied from one side

or location. When the energy is provided from multiple sites, and/or when different sources of energy are used, there is minimized risk to exceed the thermal pain tolerance. In one embodiment, the thermal energy may be applied through multiple sources, as known in the art, e.g., radiofrequency, focused ultrasound, etc. The thermal energy may be applied in a continuous manner, or it may be applied in an oscillatory manner, or it may be applied in a pulsed manner. Oscillatory or pulsed thermal energy application reduces the thermal damage to normal cells, while sufficiently heating cells to 10 which the nanoparticles are attached to the desired temperature. In this embodiment, a preferred nanoparticle is supramagnetic nanoparticle or supraparamagentic ferric oxide nanoparticle, nanotube, or nanowire.

In one embodiment, the thermal energy applied is an alter- 15 nating magnetic force applied intermittently as an alternating magnet to heat to the nanoparticle-cell complex, and an imaging system measures the temperature of the heated tissue and images the area involved. The thermal imaging system (photoacoustic or MRI) controls the energy delivery from the unit 20 to maintain the temperature at the target site at 39° C.-48° C. The thermal imaging system (photoacoustic or MRI) controls the energy delivery from the unit to maintain the temperature at the target site at 37° C.-<60° C. The thermal imaging from the unit to maintain the temperature at the target site at 37° C.-41° C. The thermal imaging system (photoacoustic or MRI) controls the energy delivery from the unit to maintain the temperature at the target site at 42° C.-46° C. The thermal imaging system (photoacoustic or MRI) controls the energy delivery from the unit to maintain the temperature at the target site at 47° C.-50° C. The thermal imaging system (photoacoustic or MRI) controls the energy delivery from the unit to maintain the temperature at the target site at 50° C.-58° C. If the nanoparticles are attached to tumor cells, the tumor cells 35 are selectively or preferentially treated over normal non-tumor cells. The thermal energy may be applied from multiple locations covering at least 1°-90°, up to 180°, and greater than 180° of the circumference of the target tissue.

The method may be used to image and treat benign or 40 malignant lesions, infective agents such as bacteria, fungi, parasites, viruses, prions, etc.

In one embodiment, the method may be used with preloaded stem cells, macrophages, dendritic cells or their exosomes to locate or track them throughout the body, imaging 45 their accumulation in vessels or tissues at a site of inflamma-

The nanoparticles may be synthetic, organic (e.g., liposomes), non-organic, non-magnetic, magnetic, paramagnetic, diamagnetic, supramagnetic, non-magnetic, mesopo- 50 rous carbide-derived carbon, iron oxide nanoparticles with gold, graphene oxide and mesoporous silicon nanostructures, carbon, quantum dots, nanoshells, nanorods, nanotubes, nanowires, quantum dots, etc. Illustrative and non-limiting specific examples also include liposomal nanoparticles, lipo- 55 some-PEG nanoparticles, micellar polymeric platform nanoparticles, L-adenine nanoparticles, L-lysine nanoparticles, PEG-deaminase nanoparticles, polycyclodextrin nanoparticles, polyglutamate nanoparticles, calcium phosphate nanoparticles, antibody-enzyme conjugated nanoparticles, poly- 60 meric lipid hybrid nanoparticles, nanoparticles containing a combination of two-three elements such as gold, gold-iron oxide, iron-zinc oxide, metallic nanoparticles, polylacticglycolic acid nanoparticles, ceramic nanoparticles, silica nanoparticles, silica crosslinked block polymer micelles, albumin- 65 based nanoparticles, albumin-PEG nanoparticles, dendrimer attached magnetic or non-magnetic nanoparticles, etc.

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Any antibodies known in the art may be used, whether targeting normal cells or tissues or pathological cells or tissues, either at a fixed static site e.g., solid tumors, or not at a fixed static site, e.g., circulating malignant cells such as blood cells, etc. Thus, the following are exemplary and not limiting: abciximab, adalimuab, alemtuzumab, brentuximab, blimubab, canakinumab, alemtuzumab, cituximab, cetrolizumab, daclizumab, denosumab, efalizumab, gemtizumab, glimumab, lipilimumab, infliximab, rituximab, muromonab, oftamumab, palivizumab, panitumab, ranibizumab, tositumomab, rastuzumab, etc.

In one specific embodiment, iron oxide nanoparticles are coated with oligosaccharides for imaging and thermotherapy. In one specific embodiment, the nanoparticles are coated with thermosensitive polymers carrying specific agents, e.g., antiinfectives, chemotherapeutics, anti-VEGFs, anti-EGFRs, hormones, immunosuppressants, immunostimulatory agents, DNA, RNA, siRNA, genes, nucleotides, etc.

In one embodiment, nanoparticles are coated or otherwise associated with biocompatible molecules. Exemplary and non-limiting biocompatible molecules include PEG, biotin, CPP, ACPP, dendrimers alone or conjugated with poly beta amine, small organic molecules, etc.

In one embodiment, enhanced cell penetration of the antisystem (photoacoustic or MRI) controls the energy delivery 25 body-nanoparticle complex may be achieved using penetration-enhancing techniques. One example of such a penetration enhancing technique is the application of ultrasound energy at the target site. One example of such a penetration enhancing technique is the use of electroporation, as is known in the art, at the target site. A combination of ultrasound and electroporation may be used. Such penetration enhancing techniques advantageously reduce the total amount of energy applied to achieve the desired effect. The result is that normal cells are less affected or not affected by the therapy, while providing the desired amount, duration, extent, etc. of therapy to the target cells.

> The combination of CPP and poly beta amine is particularly beneficial to enhance cell penetration.

> In one embodiment, enhanced immunogenicity may be achieved using agents that control the cellular immune response to DNA damage. This embodiment has utility particularly in patients undergoing cancer immunotherapy. Exemplary but not limiting agents that control the cellular immune response to DNA damage include inhibitors to CHK1, CHK2, CHM1, and CHM2. In use, such agents are attached to the nanoparticles by methods known in the art, e.g., using linkers, coatings, etc. At the target site, such agents enhance the patient's immune response to malignant cells and function synergistically with the inventive method.

> In one embodiment, cells are grown in culture with the nanoparticles. The nanoparticles are thus incorporated in the cells while in culture, and prior to injection into the body. They cells are thus associated with the nanoparticles and may be termed "nanoparticle preloaded" cultured cells. Exemplary and non-limiting cultured cell types include stem cells, macrophages, dendritic cells, lymphocytes, other leukocytes, dendritic cells, exosomes, etc. After injection of the nanoparticle preloaded cells into the body, they may be traced and imaged in various organs. Exemplary and non-limiting organs include lung, heart, brain, the respiratory system, vessels, etc. In this embodiment inflammatory process in tissues of an organ may be indirectly imaged.

In embodiment, the nanoparticles range from 1 nm-800 nm. In one embodiment, the nanoparticles range from 20 nm to 200 nm. In one embodiment, the nanoparticles range from 1 nm-20 nm. In one embodiment, the nanoparticles range from 1 nm-10 nm.

In one specific embodiment, the target site contains a lesion or pathology, and the nanoparticles contain a polymeric coating that itself contains a medication that is released when the temperature is 40° C.-47° C. In this specific embodiment, the lesion or pathology is treated by both thermally and by the 5 medication.

In one specific embodiment, the patient is treated simultaneously or with hemofiltration, hemoadsorption, mesoporous carbide-derived carbon, or another type of agent to prevent a cytokine storm.

In one embodiment, the method creates combinations of images to enhance image visibility, distinction, and utility. For example, the use of photoacoustic imaging with MRI, ultrasound, or light creates and more distinct image of a structure without heating the structure significantly beyond 15 37° C.-39° C. Similarly, while an internal structure cannot be imaged using radiofrequency wave energy or microwave energy alone, it is possible to create a photoacoustic image of an internal structure with radiofrequency wave energy or microwave energy alone if an antibody-coated nanoparticle, 20 with the antibody targeting the nanoparticle to a tumor or structure such as endothelial cells, heart cells, muscle cells, or tumor cells, etc. are initially injected, e.g. are intravenously injected. Because nanoparticles of 1 nm are significantly smaller than a micron, two nanoparticles are separated only 25 by <2 nm-3 nm, using thermal energy, they can create distinct separate signals that can be electronically enhanced. Thus, using photoacoustic technology, one can calculate and measure two points separated by that distance. This is beyond the optical resolution of a structure. The inventive method thus 30 enhances the images obtained by light, MRI, CT scan, PET scan etc.

In one embodiment, aptamers may be used in place of antibodies or monoclonal antibodies in the inventive method because of their binding specificity. Aptamers include both 35 oligonucleotides and peptides. Peptide aptamers have a short variable peptide domain, attached at both ends to a protein scaffold that can bind to various molecular targets, small molecules, proteins, nucleic acids, cells, tissues, and organisms. Aptamer may be synthesized to possess desirable storage properties and to be non-immunogenic. Aptamers may be applied for therapy of various diseases. With oligonucleotide aptamers, DNA stability is the only distinction between RNA and DNA aptamers.

Aptamers, like monoclonal antibodies, possess characteristics that make them specific for targeting. Apatamers, unlike antibodies or monoclonal antibodies, are neither thermo-sensitive nor are they damaged by heat, making them useful for thermotherapy in various applications, e.g., cancer and drug release from nanoparticles for treatment of infection, inflammatory diseases, etc. When aptamers are combined with nanoparticles and/or quantum dots, the aptamers may serve as markers for detection of tumor cells and small tumors using the inventive photoacoustic method.

In one example of this embodiment, aptamers in a composition with nanoparticles and/or quantum dots, the composition conjugated, coated, or otherwise associated with thermosensitive polymers, are used to target tumor cells for thermal energy and photoacoustic technology treatment. Thermal expansion of the complex, as a result of heat absorption, creates an internal shock wave of acoustic sound that can be recorded and imaged, i.e., photoacoustic imaging. When an aptamer-targeted thermosensitive polymer-conjugated nanoparticle and/or quantum dot complex also contains a therapeutic agent, e.g., a medicament and/or a gene or genes, 65 release of the therapeutic agent occurs when a certain temperature is achieved. The specific temperature, accurate to at

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least a tenth of a degree, is controlled using photoacoustic technology by software controlling the thermal energy delivery unit. For example, when the complex reaches a temperature of 41° C.-42° C. as a result of heating using an energy source that is under the control of a photoacoustic unit, the thermosensitive coating of the complex releases the agent, which is the medicament and/or other biomolecule and/or conjugated gene(s). Increasing the temperature of the complex, which is attached to the cell exterior or is in the cell interior, damages the cell membrane, rendering the cell, including the nucleus, more accessible to the medications, RNAi, siRNA, etc. If the temperature of the complex is increased, e.g., to 43° C.-45° C., to 47° C., up to 58° C., the cell membrane is damaged and the cell cannot survive.

Various thermosensitive polymers known in the art may be employed. For example, aptamers or variations, modifications, aptamer peptide, Affirmer, AptaBid, SELEX, etc., may be conjugated with chitosans. These chitosan-conjugated aptamer-coated nanoparticles and/or quantum dots can carry toxic medicaments, siRNA, RNAi, or genes to inhibit the uncontrolled division of, or to expedite the apoptosis process in, cancer cells. If other genes are delivered, the cells assume a different function, depending on the gene delivered.

In one example of this embodiment, thermal energy is generated externally or internally by a unit producing electromagnetic radiation, ultrasound, radiofrequency waves, microwave energy, focused ultrasound, or an alternating magnetic field. The specific temperature achieved and, in turn, the specific energy delivered, is predetermined by the operator through the photoacoustic unit that records and images the temperature of the lesion to which the nanoparticles and/or quantum dots are attached. As known to one skilled in the art, software programs are available that may superimpose or overlay the thermal image of the lesion on other images produced simultaneously by other imaging systems such as magnetic resonance imaging (MRI), magnetic resonance spectroscopy (MRS), Raman spectroscopy, ultrasound, bioluminescence, optical fluorescence, molecular Imaging (MI), imaging using contrast agents, diffusion sensitive magnetic resonance imaging, etc. This overlay ensures that both the release of the gene(s) and/or medicament by the nanoparticles and/or quantum dots, and their precise location in the body, are rendered visible in real time during the photoacoustic process.

In one example of this embodiment, the aptamer is used with a plurality of nanoparticles, quantum dots, liposomes, magnetic nanoparticles, paramagnetic nanoparticles, non-magnetic nanoparticles, synthetic nanoparticles, organic or non-organic nanoparticles, gold nanoparticles, nanowires, carbon nanotubes, graphene nanoparticles, piezoelectric nanoparticles, nanoparticles coated with chitosan, PEG, biotin, streptavidin, radionuclides, CPPs, ACCPs, etc. for cell penetration to deliver drugs or genes using photoacoustic technology and a thermal energy producing unit.

In one example of this embodiment, the antibodies or microRNAs or DNAs are obtained from a patient blood sample, and then conjugated with nanoparticles and/or quantum dots, and then administered to the patient for imaging and thermotherapy.

In one example of this embodiment, the aptamer nanoparticle and/or quantum dot complex are administered to the patient by local or systemic injection and used as a cancer bio-molecular marker for substances, whether the substances are present locally or systemically, e.g., in the circulation. The biomarkers may be detected upon removal of a blood sample and ex-vivo examination of the aptamers and/or nanoparticles and/or quantum dots. This example may also be performed in

vivo with photoacoustic technology for detection of circulating metastatic cells as described in Peyman U.S. Ser. No. 14/624,334 filed Feb. 17, 2015, or in conjunction with nanoparticles and/or quantum dots, an external energy source that heats the nanoparticles and/or quantum dots, and a photoacoustic unit as described in Peyman U.S. Patent Publication No. 2015/0087973, U.S. Ser. No. 14/554,840, filed Nov. 26, 2014, each of which is expressly incorporated by reference herein in its entirety,

In one example of this embodiment, a biomarker such as microRNA, DNA, proteins, etc. is obtained from the blood of a patient prior to mechanical manipulation of tissue in which the tumor may reside, e.g., prostate, breast, liver, spleen, colon, eye, etc., and compared with post-manipulation values using low impact trauma, massage, shaking, low frequency ultrasound, or mechanical vibration of an organ. The biomarkers obtained are then conjugated with aptamers and/or nanoparticles and/or quantum dots for thermotherapy and imaging.

In one example of this embodiment, the aptamer and nanoparticle and/or quantum dot complex is also conjugated with immune stimulating agents and the method is performed. In this example, cancer cells are destroyed with locally applied thermal energy through the nanoparticles, and additionally immune cells are stimulated in order to remove the damaged or non-damaged cells.

In one example of this embodiment, aptamer and nanoparticle and/or quantum dot complex is conjugated with at least one radionuclide to effect cell destruction by both local radiation and hyperthermal therapy. In this example, the vascular supply to the tumor site is also damaged by the combination of targeted hyperthermal therapy with localized radiotherapy. Radionuclides and methods of radiotherapy are known to one skilled in the art; radionuclides include but are not limited to 35 and 32 P.

In one example of this embodiment, the aptamer and nanoparticle and/or quantum dot complex is administered locally or systemically to discover and localize small tumors prior to progression of a fully-progressed metastatic process. Attachment of the aptamers and/or nanoparticles and/or quantum dots to byproducts of cancer can be used as a targeting molecule to photoacoustically image the cancer cells. In small-sized tumors, this example is preferably used in combination with other imaging technologies such as MRI, positron, CT-scan, ultrasound, OCT, etc. to treat the lesion simultaneously with thermotherapy and localized drug release.

In one example of this embodiment, a plurality of aptamers conjugated with nanoparticles and/or quantum dots are used in addition to antibodies (polyclonal or monoclonal) that are 50 also conjugated with nanoparticles and/or quantum dots, to achieve a broader thermal effect than may be achieved using hyperthemal treatment. For example, the pegylated antivascular growth factor aptamer pegaptinib (Macugen®, Pfizer) was not very successful in treating the wet form of age related 55 macular degeneration. This is because Macugen® blocked only a specific vascular endothelial growth factor (VEGF) receptor in ocular tissue. In contrast, bevacizumab (Avastin®, Genentech/Roche) and ranibizumab (Lucentis®, Genentech) bound to a wider variety of VEGF receptors and were more 60 efficacious but, as a major drawback, required repeated intraocular injections resulting in patient discomfort. However, the combination of nanoparticles and/or quantum dots/ and aptamer and/or antibodies, as described herein, provide therapy by damaging endothelial cells of the new vessels with thermal energy, while an anti-VEGF agent incorporated in the nanoparticles and/or quantum dot complex enhances the

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effect of the therapeutic medication on the abnormal vessels, and entirely eliminates the need for repeated intraocular injection

In one example of this embodiment, the aptamer-conjugated nanoparticle and/or quantum dot complex is conjugated with one or more thermosensentive polymer(s) and injected either in the vitreous cavity or in the circulation. Within a relatively short time thereafter, the area of neovascularization is treated with low-level laser energy, under an operator's control of the photoacoustic technology, to release the anti-VEGF medication and to thermally damage the abnormallygrowing endothelial cells and/or vessels by creating a vascular occlusion, thus preventing bleeding and regrowth of new vessels. In one embodiment, intraocular tumors such as melanomas or retinoblastomas are treated with aptamers and/or nanoparticles and/or quantum dots conjugated with the antitumor agent. In this embodiment the thermal energy can be delivered as focused ultrasound, radiofrequency waves, microwaves, or using an alternating magnet and magnetic or 20 paramagnetic nanoparticles. Tumors in the body, irrespective of location and not limited to the brain, are treated in the same manner.

In one example of this embodiment, the aptamer and nanoparticle and/or quantum dot complex is additionally conjugated with a photosensitizer that, upon laser application, is activated to result in more permanent damage to the malignant cells. Because malignant cells in the body constantly die and multiply simultaneously, their microDNA or DNA particles circulate in the serum before the tumor has achieved a size of about 5 mm³, the size at which tumors typically metastasize and/or can be visualized by imaging technology other than the inventive photoacoustic technology.

In one example of this embodiment, aptamers and/or nanoparticles and/or quantum dots are conjugated with antibodies to obtain enhanced tumor cell recognition from cancer cell biomolecules present in the circulation. These tumors can then be targeted with thermotherapy using photoacoustic technology and using the antibody or aptamer as a biomarker for imaging and thermal delivery of the aptamer and/or nanoparticle and/or quantum dot to the lesion.

In one example of this embodiment, the aptamer and nanoparticle and/or quantum dot complex is coated with an appropriate antibacterial, antifungal, antiviral, anti-parasitic surface antigen directed toward the target bacteria, fungus, virus, or parasite, respectively. When injected locally or systemically, these aptamers and/or nanoparticles and/or quantum dots are attached to the targets, where they are heated using thermal energy, controlled by photoacoustic technology, to release the medication and simultaneously heat the cell membrane of the bacteria, fungus, virus, or parasite to 45° C.-47° C. or higher to damage and eliminate them.

In one example of this embodiment, the aptamer and nanoparticle and/or quantum dot complex is conjugated or otherwise associate with antibodies against beta-amyloid and toxic Tau protein that is present in Alzheimer's disease. The conjugates are then injected, e.g., into cerebrospinal fluid, locally, or systemically, to release medication against beta-amyloid and Tau protein with thermotherapy. Heating the beta amyloid or Tau protein also stimulates microglial migration and thus removes the remaining debris.

There are numerous indications for gene and/or drug therapy using the inventive method, i.e., temperature sensitive polymers conjugated with aptamers and/or nanoparticles and/or quantum dots. One benefits is its specificity to a particular organ, tissue, or even cell; it is a targeted process. Another benefit is that release of the gene and/or medicament is stringently controlled by the operator, and occurs only at a

pre-defined specific temperature. The result is controlled, local release and elimination of systemic side effects of the therapeutic agents.

In one example of this embodiment, an injured spinal cord nerve or peripheral nerve is treated. A opsin family gene is 5 provided with the aptamers and/or nanoparticles and/or quantum dots for subsequent therapeutic light pulse stimulation to assist in recovery of damaged nerves and eliminating viral transfection that produces hypersensitivity. For example, in one embodiment an opsin family gene is delivered to the 10 desired cells for subsequent stimulation with visible or infrared light. In another embodiment, the missing gene(s) in an organism are replaced using the aptamers and/or nanoparticles and/or quantum dots containing the gene(s), which are released by increasing the temperature of the aptamers and/or 15 nanoparticles and/or quantum dots. The release is monitoring using photoacoustic technology which indicates achievement of the specific release temperature. The thermal energy can be electromagnetic radiation, microwave radiation, radiofrequency waves, ultrasound, etc. As an example, an opsin gene 20 family can be administered in the eye, in the vitreous cavity, or under the retina to reach brain cells, or by local injection into the cerebrospinal fluid or over the brain, or for selective stimulation of neurons in stroke, epilepsy, depression, Parkinson's disease, etc. Subsequent stimulation with appropri- 25 ate light pulses can be done deliberately by the patient or by a processor, controlling the pulse duration, etc. to achieve the desired therapeutic effect. This effect may be, e.g., object recognition by a degenerative retina, epilepsy cessation in a stroke patient, alleviating depression in a patient with posttraumatic brain syndrome, or regaining function in spinal cord or peripheral nerve injury.

In one example of this embodiment, a composition of the aptamer and nanoparticle and/or quantum dot complex con- 35 taining opsin family genes is administered to excitable cells, e.g., retinal cells in degenerative diseases such as retinitis pigmentosa, brain cells in Alzheimer's or Parkinson's disease, nerve cells in spinal cord injury, or cardiac cells in arrhythmia or myocardial infarction. The gene-transfected 40 cells are then stimulated by light, inducing an action potential in the membrane of the excitable cells. By replacing the missing genes, there is a potential to reverse the disease process.

In one example of this embodiment, local injection of the 45 aptamer and nanoparticle and/or quantum dot complex is preferred over systemic injection. Local injection may be, e.g., in the eye, in the central nervous system, in the cerebrospinal fluid, etc. Local injection at a site within the blood brain barrier prevents contact of the composition with plasma 50 enzymes that can damage the DNA carried by the composition entering these areas. Alternatively, or when local administration is not feasible, systemic administration of the aptamer and/or nanoparticle and/or quantum dot-gene complex is performed with the complex containing a PEG coating 55 of sufficient thickness to minimize or prevent damage by plasma enzymes to DNA and/or RNA. After administration, the aptamer and nanoparticle and/or quantum dot complex enters the specific cells via receptor mediated uptake. Once internalized, the DNA and/or RNA is released from the cyto- 60 plasmic endosomes or passes through nuclear pores. In the nucleus, the opsin gene or other genes are released, replacing the missing or the defective gene. For example, Parkinson's disease can be treated with the opsin gene family and light pulses used to stimulate function of the remaining neurons 65 and also to control involuntary motion of these patients. Thermally induced opsin genes or other genes may be delivered to

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the retina, brain, spinal cord, cardiac cells, and/or peripheral nerves to subsequently stimulate these cells with either an internally implanted fiber optic, or an externally delivered series of light pulses. The resulting therapeutic effect is, e.g., regulated cardia rhythm, ameliorating epilepsy, brain stimulation replacing existing electrical stimulation requiring wires that induce scar formation and require periodic replacement. As another example, patients with retinal genetic diseases such as retinitis pigmentosa, Leber amaurosis congenita, and retinoschisis may be treated by the inventive method where the opsin gene family is provided along with the gene that is lacking in these patients.

In one embodiment dopamine or opsin genes are conjugated with targeted nanoparticles and are delivered through the nose by spraying, drops, or injection. After accessing the olfactory nerves, the nanoparticles readily travel to various areas in the brain. One can subsequently stimulate different brain areas with light applied externally to or through the nose, e.g., to control involuntary tremors in Parkinson's disease, ameliorate Alzheimer's disease, control epilepsy, encourage functionalization of the brain after cerebral vascular infarct, etc. For example, piezoelectric nanoparticles, previously described, may be delivered to the hose, and brain stimulation can be performed by an external ultrasound applied over the nose.

### **EXAMPLE**

FIG. 2 illustrates a patient with a choroidal tumor, diagbrain after a trauma, assisting recovery of loss of a function in 30 nosed clinically as malignant melanoma. The patient is treated with the inventive thermotherapy using ferromagnetic nanoparticles coated with (poly)ethylene glycol, anti-tumor antibody, and a polymer having anti-cancer medication that releases the medication when the temperature reaches 42° C. The ferromagnetic nanoparticles are also coated with radioactive molecules that emit  $\alpha$  or  $\beta$  radiation with a short halflife (e.g., 3 hours to days). The nanoparticles are administered intravenously to the patient. A natural magnet or an electromagnet is sutured to the sclera if the tumor cannot be covered with the magnet if placed externally as shown in FIG. 2. If the tumor is located anteriorly, an electromagnet is preferably applied non-invasively, through the conjunctiva to the tumor as shown in FIG. 3. After a waiting period, e.g., one minute to one hour, sufficient additional ferromagnetic particles are accumulated in the tumor. After the magnet is removed, the patient is treated locally by the same electromagnet using a rapidly alternating magnetic field. However, when it is desirable to treat the tumor and possible metastatic lesions, a large reversible magnet covering the entire body is used and the patient's entire body receives thermotherapy.

By rapidly reversing the magnetic field, the ferromagnetic nanoparticles are heated due to an internal semicircular motion in the individual ferromagnetic nanoparticles. The degree of heating is recorded by the photoacoustic system which, in turn, is communicated by a processor with the reversible magnet to generate the desired temperature at the tumor site. When the desired temperature is achieved, this desired temperature is maintained for a desired period, e.g., about 15 minutes in one embodiment, greater than 15 minutes in another embodiment. In general, the initial thermal energy is delivered to achieve a temperature of 42° C. to 43° C. to release the drugs from the nanoparticles. If needed, the temperature is gradually increased to, e.g., 45° C., 47° C., 50° C., or greater than 50° C. for a period, as needed. Parameters such as these are programmed by processor software that controls thermal energy delivery. FIG. 6 illustrates how information obtained from the photoacoustic system controls the energy

delivery by the processor to achieve and maintain the desired temperature, for a desired time, in the nanoparticle-cell complex at the tumor or lesion site.

After thermotherapy is completed, the patient is administered topical antibiotics for a few days and remain under physician observation. The treatment may be repeated at intervals of 3 months to 9 months, or more or less frequently as needed, to destroy the tumor. The release of thermal energy, radioactivity, and drugs during the therapy generally gradually damages tumor cells and permits gradual patient recovery. Because the therapy is generally not invasive, or is only minimally invasive in embodiments where an internal magnet is used, the patient does not require long-term hospitalization. In a patient with severe disease, the patient may be treated with a low dose of anti-cancer medication to further kill the already damaged tumor cells, without damaging healthy cells.

In one embodiment after placement of the magnet, the ferromagnetic nanoparticles, which may additionally be 20 radioactive and/or contain a therapeutic agent, are concentrated at a benign or malignant tumor after systemic or local administration at a localized site, e.g., in the eye, skin, breast, extremities, CNS, spinal cord, vertebra, pelvis, urogenital system, tongue, throat, etc. Examples are shown in FIGS. 4A, 25 4B for a tumor affecting the lower leg and with a horseshoe-shaped electromagnet configuration, and in FIG. 5 for a facial skin tumor.

In one embodiment, the treatment of the tumor requires medication only. In this embodiment, the ferromagnetic 30 nanoparticles carry medication in the absence of a radioactive molecule. The ferromagnetic nanoparticles may carry, e.g., immune modulators, anti-VEGF, sRNA, RNA, genes, bacteriophages, antibiotics, growth hormones, growth inhibitors, and combinations of these as desired.

The references disclosed and cited below are expressly incorporated by reference herein in their entirety.

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Although several embodiments have been chosen to illustrate the invention, those skilled in the art will readily appreciate that various changes and modifications can be made without departing from the scope of the invention.

## What is claimed is:

1. A method using hyperthermal therapy for at least one of 10 delivery of a therapeutic agent to a cell in a patient in need thereof, the method comprising the steps of:

providing a targeted nanoparticle-agent to form a nanoparticle-agent-cell complex at a site,

thereafter exposing the site to an energy source under conditions sufficient to achieve a temperature of the complex, for about 1 femtosecond to about 15 minutes, the temperature sufficient result in agent release from the complex to the cell by at least one of

rupture of cellular membranes of cells in contact with or 20 proximate to the nanoparticles, or

formation of holes or pores in the cellular membrane; thereafter assessing the acoustic sound produced from the nanoparticle at the site and,

correlating the acoustic sound with the presence or absence 25 of viable cells at the site by photoacoustic imaging and/ or ultrasound spectroscopy.

- 2. The method of claim 1 where the therapeutic agent is at least one of a gene or a non-gene therapeutic agent, and where the agent is a gene the gene replaces or corrects a genetic 30 defect in the cell.
- 3. The method of claim 2 where the therapy is applied to a patient with cancer.
- **4.** The method of claim **3** where the agent is an anti-cancer agent, a tumor suppressor gene, and combinations thereof.
- 5. The method of claim 1 where the hyperthermal therapy is applied to the patient for general or en masse treatment of an organ or body region.
- **6**. The method of claim **1** where hyperthermal therapy is applied for a duration between one femtosecond to one second.
- 7. The method of claim 1 where hyperthermal therapy achieves a temperature of at least  $60^{\circ}$  C.
- 8. The method of claim 1 further comprising administering nanoparticles coated with a thermosensitive polymer to the 45 patient to verify that the nanoparticles are in contact with the appropriate tissue, including circulating or non-circulating cells, and using a photoacoustic unit to achieve a temperature of 37° C. to 43° C. to release the agent attached to the nanoparticles.
- 9. The method of claim 1 resulting in damaged cells or damaged exosomes, and further comprising quantitating at least one of the resultant damaged cells or damaged exosomes to assess efficacy.
- 10. The method of claim 9 where quantitating is by a 55 patient wrist band counter or by a patient neck collar counter.
- 11. The method of claim 1 where a complex of a piezoelectric nanoparticle and a gene is administered to a patient, and an externally-positioned ultrasound source activates the complex to control cell polarization, the method capable of complex activation in the absence of light penetration.
- 12. The method of claim 1 where the nanoparticles are selected among a plurality of quantum dots, hybrid quantum dots, non-quantum dot nanoparticles, nanoparticle types, nanoparticle components, nanoparticle complexes, nanowires, nanoshells, nanorods, caged nanoparticles, nanocrystals, nanotubes, nanobots, and nanoparticle compositions, non-

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magnetic, magnetic, paramagnetic, diamagnetic, supramagnetic, non-magnetic, piezoelectric, liposomes, dendrimers, and combinations thereof.

- 13. The method of claim 1 where the nanoparticles comprise compositions selected from the group consisting of organic, non-organic, metallic, non-metallic, silver, lanthanide, cerium, gold, zinc, silicon, perovskite, ceramic, plastic, gallium/arsenide, cadmium/selenium, copper/indium/ gallium/selenide, zinc sulfide, indium/gallium/phosphide, gallium arsenide, indium/gallium nitride, a nanocrystal and a metal, mesoporous carbide-derived carbon, iron oxide nanoparticles with gold, graphene oxide, mesoporous silicon nanostructures, particles of semiconductor/metal and semiconductor/semiconductor hetero-junctions, biocompatible short peptides of naturally occurring amino acids that have the optical and electronic properties of semiconductor nanocrystals, graphene quantum dots, graphene-oxide quantum dots, graphene-zinc oxide quantum dots, graphene-zinc oxide quantum nanowires, carbon nanotubes, carbon-based skeletal-type nanostructures, fullerenes, bucky balls, micelles, micelle-like structures, liposomes, dendrimers, aptamers, single-stranded deoxyribonucleic acid- or ribonucleic acidoligonucleotide aptamer-conjugated or modified nanoparticles, dendrimers including poly(amidoamine) (PAMAM), poly(amidoamine-organosilicon) (PAMAMOS), poly(propyleneimine) (PPIO), tecto, multilingual, chiral, hybrid, amphiphilic, Frechet-type dendrimers, liposome-PEG nanoparticles, micellar polymeric platform nanoparticles, L-adenine nanoparticles, L-lysine nanoparticles, PEG-deaminase nanoparticles, polycyclodextrin nanoparticles, glutamate nanoparticles, calcium phosphate nanoparticles, antibody-enzyme conjugated nanoparticles, polymeric lipid hybrid nanoparticles, nanoparticles containing a combination of two-three elements such as gold, gold-iron oxide, iron-zinc oxide, metallic nanoparticles, polylacticglycolic acid nanoparticles, ceramic nanoparticles, plasmonic nanoparticles, silica nanoparticles, silica crosslinked block polymer micelles, albumin-based nanoparticles, albumin-PEG nanoparticles, dendrimer attached magnetic or non-magnetic nanoparticles, graphene/zinc oxide (ZnO) and reduced graphene oxide, plasmonic nanoparticles coated with reduced graphene oxide, dextran-reduced graphene oxide, and combinations thereof.
- 14. The method of claim 1 where the agent is at least one gene selected from the group consisting of genes involved in cell membrane polarization and depolarization, genes encoding opsin family genes, genes encoding neurotransmitters, genes encoding regulatory proteins, genes encoding products for synthesis or inhibition of a therapeutic protein, tumor suppressor genes, therapeutic genes, defective genes, genes that have an stimulatory action in response to light or ultrasound, and combinations thereof.
- 15. The method of claim 1 where the energy types to activate the nanoparticles is selected from the group consisting of electromagnetic radiation, visible light, invisible light, alternating magnet, reversible magnet, and combinations thereof.
- 16. The method of claim 1 further comprising administering a clustered regularly interspersed palindromic repeats (CRISPR) with the nanoparticles for gene repair, gene editing, and combinations thereof.
- 17. The method of claim 1 further comprising providing a nanoparticle associated with a member of specific binding pair, a cellular receptor-agonist or antagonist pair, an enzyme-substrate pair, and/or an antibody-antigen pair, a virus, or virus like particles (VLP).

- 18. The method of claim 1 further comprising providing a nanoparticle associated with a virus like particle resulting in an enhanced immune therapy to a tumor cell antigen in the absence of an adjuvant.
- **19**. The method of claim **1** further comprising providing a nanoparticle with a biocompatability enhancing coating.
- 20. The method of claim 19 where the coating is a biocompatible polymer selected from the group consisting of biotin, streptavidin, (poly)ethylene glycol, chitosan, cell penetration peptide (CPP), arginine CPP, and combinations thereof.
- 21. The method of claim 1 where the nanoparticle size ranges from 1 nm to greater than 999 nm.
- 22. The method of claim 1 where the gene is an opsin family gene, and the energy frequency provided is selected from the group consisting of
  - >30 Hz with repeated pulses resulting in decreased cellular survival due to enhanced cell vulnerability to the agent, and

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- <20 Hz to 30 Hz with repeated pulses resulting in enhanced cellular survival or normal cell multiplication in the presence of growth factors.
- 23. A method using hyperthermal therapy for at least one of delivery of an therapeutic agent to a cultured cell, the method comprising the steps of:
  - providing a nanoparticle containing or complexed to an agent to form a nanoparticle-agent-cell complex,
  - thereafter exposing the site to an energy source under conditions sufficient to achieve a temperature of the complex sufficient result release of the agent from the complex to the cells by at least one of
    - rupture of cellular membranes of cells in contact with or proximate to the nanoparticles, or

formation of holes or pores in the cellular membrane.

\* \* \* \* \*

## UNITED STATES PATENT AND TRADEMARK OFFICE

## CERTIFICATE OF CORRECTION

PATENT NO. : 9,393,396 B2 Page 1 of 2

APPLICATION NO. : 14/869051
DATED : July 19, 2016
INVENTOR(S) : Peyman

It is certified that error appears in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

### TITLE PAGE

Column 1, (63) Related U.S. Application Data currently reads:

"Continuation of application No. 14/724,577, filed on Apr. 6, 2015, now abandoned, which is a" Column 1, (63) Related U.S. Application Data should read:

"Continuation of application No. 14/724,577, filed on May 28, 2015, now Pat No. 9,320,813, which is a"

Page 2, Column 1, Related U.S. Application Data currently reads:

"continuation-in-part of application No. 14/679,083, filed on Feb. 17, 2015, now Pat. No. 9,320,813, which is a continuation-in-part of application No. 14/624,334, filed on Feb. 17, 2015, now Pat. No. 9,302,087, which is a continuation-in-part of application No. 14/554,840, filed on Nov. 26, 2014, now Pat. No. 9,017,729, which is a continuation-in-part of application No. 14/311,464, filed on Jun. 23, 2014, now Pat. No. 8,932,636, which is a continuation-in-part of application No. 13/915,282, filed on Jun. 11, 2013, now Pat. No. 8,668,935, which is a continuation-in-part of application No. 13/665,458, filed on Oct. 31, 2012, now Pat. No. 8,709,488, which is a continuation-in-part of application No. 13/610,503, filed on Sep. 11, 2012, now Pat. No. 8,795,251, which is a continuation-in-part of application No. 13/527,005, filed on Jun. 19,2012, now Pat. No. 8,808,268, which is a continuation-inpart of application No. 13/455,237, filed on Apr. 25, 2012, now Pat. No. 8,801,690, which is a continuation-in-part of application No. 13/361,786, filed on Jan. 30, 2012, now Pat. No. 8,481,082, which is a continuation-in-part of application No. 13/307,916, filed on Nov. 30, 2011, now Pat. No. 8,119,165, which is a continuation-in-part of application No. 13/189,606, filed on Jul. 25, 2011, now Pat. No. 8,137,698, which is a continuation-in-part of application No. 13/149,209, filed on May 31, 2011, now Pat. No. 7,964,214, which is a continuation-in-part of application No. 12/478,029, filed on Jun. 4, 2009, now Pat. No. 7,638,139, which is a continuation-in-part of application No. 11/485,352, filed on Jul. 13, 2006, now Pat. No. 7,101,571."

Page 2, Column 1, Related U.S. Application Data should read:

"continuation-in-part of application No. 14/679,083, filed on Apr. 6, 2015, now Pat. No. 9,302,087, which is a continuation-in-part of application No. 14/624,334, filed on Feb. 17, 2015, now Pat. No. 9,289,491, which is a continuation-in-part of application No. 14/554,840, filed on Nov. 26, 2014, now Pat. No. 9,233,157, which is a continuation-in-part of application No. 14/311,464, filed on Jun. 23,

Signed and Sealed this Twenty-seventh Day of September, 2016

Michelle K. Lee

Michelle K. Lee

Director of the United States Patent and Trademark Office

# CERTIFICATE OF CORRECTION (continued) U.S. Pat. No. 9,393,396 B2

2014, now Pat. No. 9,017,729, which is a continuation-in-part of application No. 13/915,282, filed on Jun. 11, 2013, now Pat. No. 8,932,636, which is a continuation-in-part of application No. 13/665,458, filed on Oct. 31, 2012, now Pat. No. 8,668,935, which is a continuation-in-part of application No. 13/610,503, filed on Sep. 11, 2012, now Pat. No. 8,709,488, which is a continuation-in-part of application No. 13/527,005, filed on Jun. 19, 2012, now Pat. No. 8,795,251, which is a continuation-in-part of application No. 13/455,237, filed on Apr. 25, 2012, now Pat. No. 8,808,268, which is a continuation-in-part of application No. 13/361,786, filed on Jan. 30, 2012, now Pat. No. 8,801,690, which is a continuation-in-part of application No. 13/307,916, filed on Nov. 30, 2011, now Pat. No. 8,481,082, which is a continuation-in-part of application No. 13/189,606, filed on Jul. 25, 2011, now Pat. No. 8,119,165, which is a continuation-in-part of application No. 13/149,209, filed on May 31, 2011, now Pat. No. 8,137,698, which is a continuation-in-part of application No. 12/478,029, filed on Jun. 4, 2009, now Pat. No. 7,964,214, which is a continuation-in-part of application No. 11/485,352, filed on Jul. 13, 2006, now Pat. No. 7,638,139, which is a division of application No. 10/073,863, filed on Feb. 14, 2002, now Pat. No. 7,101,571."